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(54) Title: POLYPEPTIDE FOR REPAIRING GENETIC INFORMATION, NUCLEOTIDIC SEQUENCE WHICH CODES FOR IT AND PROCESS FOR THE PREPARATION THEREOF (GUANINE THYMINE BINDING PROTEIN - GTBP)

(57) Abstract

The present invention relates to a new protein, GTBP (Guanine Thymine Binding Protein), that binds to G/T DNA mismatches to mediate repair of genetic information, to methods for detection of this protein, to the nucleotidic sequence encoding this protein and to processes for obtaining the abovementioned protein using genetic engineering techniques. Furthermore, the present invention has as its object the detection in tumor tissues of the mutant GTBP gene in order to prevent and provide rapid diagnosis of human colorectal tumor forms. The figure shows the absence of GTBP-specific activity in cells obtained from human colorectal tumors.

HeLa LoVo DLD1 G/C G/T G/C G/T G/C G/T

specific complex

non-specific complexes



←free probe

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POLYPEPTIDE FOR REPAIRING GENETIC INFORMATION, NUCLEOTIDIC SEQUENCE WHICH CODES FOR IT AND PROCESS FOR THE PREPARATION THEREOF (GUANINE THYMINE BINDING PROTEIN - GTBP).

DESCRIPTION

Technical field

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This invention relates to the area of cancer prevention, diagnosis and therapeutics. In particular, the invention is concerned with methods for detection of a novel mismatch binding protein, termed GTBP (Guanine Timine Binding Protein), which mediates the repair of genetic information, with the nucleic acid encoding the protein and with processes for obtaining the and producing it by recombinant protein techniques. In addition, engineering the present invention also relates to detection of mutated GTBP gene in tumour tissues and to prevention and early diagnosis of human colorectal cancers.

Background of the discovery

In human cells, mismatch recognition and binding has until now been believed to be mediated by the hMSH2 protein. The observation that cells from human colorectal cancers (CRC) exhibit a mutator phenotype with a marked instability of microsatellite sequences suggested that these tumor cells may be deficient in DNA mismatch repair. This hypothesis was substantiated when extracts from CRC tumor-derived cell lines where shown to be unable to repair mismatches in an in vitro assay (see refs. 1 and 2 for reviews).

The serendipitous discovery of an open reading frame (ORF) encoding a polypeptide homolog of the E. coli mismatch-binding protein MutS (3, 4) paved the way for the identification of an ever-growing family of MSH genes, ranging from bacteria to man (see e.g. 5). Three members of this family, S. cerevisiae MutS homologs MSHl and MSH2, as well as the human homolog hMSH2, could be shown to bind to mismatched DNA in vitro (6-9). The link between the biological function of hMSH2 and the

phenotype of the CRC tumors was forged when (i) the hMSH2 gene was shown to segregate with a known CRC locus on chromosome 2p (10,11), (ii) the hMSH2-deficient cell line LoVo was shown to be deficient in mismatch repair (12) as 5 well as in mismatch-binding activity (13) and (iii) the genome of this cell line exhibited a marked instability microsatollite sequences (14). A mismatch-binding (for <u>G/T</u> binding protein), originally identified in HeLa cells by the present inventors (15), 10 shown to bind preferentially to heteroduplexes containing G/T mispairs. Purification of this DNA binding activity by G/T mismatch affinity chromatography yielded a mixture of two polypeptides of apparent molecular weights of 100 and 160 kDa (16), indicating that the mismatch-specific complex was composed of two proteins. 15 The 100 kDa constituent of the complex was demonstrated to be hMSH2 (17). The present discovery implies that hMSH2 acts as a complex with GTBP in the correction of base/base mispairs and one- or two-nucleotide loops. Moreover, GTBP is necessary but not indispensible in the 20 correction of larger insertion/deletion loops. A number of tumors have been shown to display mutator phenotypes which are consistent with the functional role of the hMSH2-GTBP complex (20-24). Prior the to discovery and characterization of GTBP, no specific role 25 in the repair of genetic information and no hereditary defect had been associated with this protein or with the gene encoding it.

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30 Sumary of the invention

It is an object of the present invention to provide a 1360-amino acid sequence corresponding to the polypeptide referred to as GTBP. It should be stated that GTBP is used to indicate a compound polypeptide combining in order the amino acid sequences indicated in SEQ ID NO:15 (from amino acid 1 to 68) and SEQ ID NO:1 (from amino acid 1 to 1292).

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It is another object of the present invention to provide a genetic construct containing a double-stranded cDNA sequence of 4080 base pairs encoding a 1360-amino acid peptide referred to as GTBP. It should be stated that the whole coding gene GTBP indicates a compound DNA sequence combining in order the nucleotide sequences indicated in SEQ ID NO:16 (from nucleotide 1 to 204) and SEQ ID NO:12 (from nucleotide 1 to 3980).

A further object of the present invention is to provide a genetic construct capable of expressing a 1360-amino acid peptide of molecular mass 153 kDa referred to as GTBP.

It is another object of the present invention to provide a method for preparation and isolation of native GTBP protein in pure form from cultured cells and tissues.

It is another object of the present invention to provide a method for the assessment of the *in vitro* activity of GTBP.

It is yet another object of the present invention to provide a method for the detection of mutated GTBP by the use of specific antibodies directed against GTBP.

It is yet another object of the present invention to provide a method for the detection of mutated *GTBP* alleles by the use of the polymerase chain reaction and sequencing of the amplification products.

It is another object of the present invention to provide DNA probes for the detection of mutated GTBP genes in human cells.

It is an object of the present invention to provide a method for diagnosing and prognosing of human colorectal cancers (CRC).

It is yet another object of the present invention to provide a method for detecting the genetic predisposition to human colorectal cancers (CRC).

It is yet another object of the present invention to provide a method for large-scale population screening to genetic predisposition to human colorectal cancers (CRC).

It is still another object of the present invention to provide a method for supplying wild-type *GTBP* alleles to a cell which has lost the *GTBP* gene function.

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It is another object of the present invention to provide a method for generating transgenic animals carrying mutant GTBP alleles.

It is another object of the present invention to provide a method for testing the activity of therapeutic agents aimed to suppress human colorectal cancers (CRC).

These and other objects of the invention are provided by one or more of the embodiements which are described below.

In one embodiment the sequence of a 1360-amino acid polypeptide is provided corresponding to the protein referred to as GTBP.

In another embodiment a cDNA molecule is provided which comprises the coding sequence of the GTBP gene.

In another embodiment a procedure for the preparation of the pure GTBP protein is provided.

It is another embodiment of the present invention to provide pairs of single stranded primers to determine the nucleotide sequence of the *GTBP* gene or of DNA regions internal to the *GTBP* gene by polymerase chain reaction. The sequence of said primers is internal to chromosome 2p16, said pairs of primers allowing the syntesis of *GTBP* gene or of parts of it.

In yet another embodiment of the present invention a nucleic acid probe is provided which is complementary to human wild-type GTBP gene coding sequence and which can form mismatches when annealed with mutant GTBP alleles, thereby making possible the detection of heteroduplex DNA as revealed by shifts in electrophoretic mobility either with or without prior enzymatic or chemical cleavage.

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In another embodiment a procedure is indicated for the detection of wild-type or mutated GTBP protein in humans, comprising: isolating a human sample selected from the tissue or body fluid and detecting the wild-type or the altered GTBP protein itself or in any complex formed by the association of GTBP with other polypeptides.

In another embodiment of the present invention a method is provided for the assessment of the activitiy of (i) the wild-type GTBP protein or (ii) of derived peptides obtained by deletion or insertion of known amino acid sequences in GTBP protein or (iii) of the altered GTBP protein as the result of in vivo mutational events or (iv) of any complex formed by the association of peptides just mentioned in (i), (ii), (iii), and (iv) of the present embodiment with other polypeptides.

In yet another embodiment a method is provided for the detection of cancer in humans, comprising: isolating a human sample selected from the tissue or body fluid; detecting the alteration in the GTBP gene or in the expressed polypeptide (GTBP protein) itself or in any complex formed by the association of GTBP with other polypeptides, said alteration indicating the predisposition to neoplastic transformation or the presence of cancer.

In still another embodiment of the present invention a method of diagnosing or prognosing neoplastic tissue of a human is provided comprising: detecting somatic alterations in wild-type GTBP alleles or their expression products in human colorectal cancers (CRC), said alteration indicating neoplasia of the tissue.

In yet another embodiment a method is provided for the detection of genetic predisposition to CRC, comprising: isolating a human sample selected from the group consisting of blood, bioptic samples of tissues, esfoliative cells and any other generic human sample; detecting the alteration in the GTBP gene or in the

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expressed polypeptide (GTBP protein) itself or in any complex formed by the association of GTBP with other polypeptides, said alteration indicating genetic predisposition to cancer.

In another embodiment of the present invention a method is provided for supplying wild-type GTBP gene function to a cell which has lost said gene function by virtue of any mutation in the GTBP gene, comprising: introducing wild type GTBP gene into a cell which has lost said gene function such that GTBP gene is then expressed at wild-type level in the cell. GTBP protein can also be applied to cells or administered to animals to remediate defects in GTBP gene function.

In an additional embodiment a method is provided to supply a portion of wild-type GTBP gene to a cell which has lost the said gene such that the said portion is expressed in the cells and encodes part of the GTBP protein which is required for non-neoplastic growth of the said cell.

It is another embodiment of the present invention the generation of transgenic animals carrying a mutated GTBP gene derived from a second species or a mutated GTBP gene generated in vitro by genetic engineering techniques.

In another embodiment of the present invention a method of testing therapeutic agents for the ability to suppress a neoplastically trasformed phenotype is provided. The method comprises: applying a test substance to a cultured epithelial cell which carries a mutation of the GTBP gene and determining whether the substance suppresses the neoplastic phenotype of the cell or suppresses the growth of already developed tumors.

In another embodiment of the present invention a method of testing therapeutic agents for the ability to suppress a neoplastically trasformed phenotype is provided. The method comprises: applying a test substance to an animal which carries a mutation of the GTBP gene

and determining whether the substance prevents neoplastic transformation of defined tissues or suppresses the growth of already developed tumors.

The present information provides the art with the information that the GTBP gene, a heretofore unknown gene, encodes the GTBP protein which acts as specific mismatch-binding factor. GTBP binds preferentially to heteroduplexes containing G/T mispairs and one- or two-Purification of this nucleotide loops. DNA activity made it possible to establish that the mismatchspecific factor is in fact a complex composed of two distinct proteins. The smaller constituent of the complex (about 100 kDa) is the hMSH2 protein (17) whereas the larger component (about 160 kDa) is GTBP. The present invention provides the technical tools for the detection and for the activity assessment of GTBP alone or as a complex with hMSH2. The GTBP gene is a target mutational events, these alterations being associated with tumorigenesis. This discovery allows highly specific assays to be performed to determine the neoplastic status of a particular tissue or the predisposition to cancer of individuals. A number of tumors have been shown to display mutator phenotypes with a similarly low degree of microsatellite instability (20-24) consistent with the functional role of the hMSH2-GTBP complex. Prior to the discovery and characterization of current GTBP. specific role in the repair of genetic information and no hereditary defect had been associated with this protein.

Brief description of the drawings.

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Figure 1 a shows the commercial phagemid vector pBluescript SK⁻ (Stratagene) used for cloning and sequencing the GTBP cDNA. The DNA fragment shown in SEQ ID NO: 12 was cloned between the *EcoRI* and *XhoI* sites of the vector. b shows the commercial pCITE 2b vector. The insert described in SEQ ID NO: 12 was inserted between the *EcoRI* and *XhoI* sites of the vector.

Ampicillin = beta-lactamase gene for ampicillin resistance

ColE1 ori = origin of replication derived from plasmid
ColE1

5 f1 = origin of replication of phage F1

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lacZ = alpha peptide of beta-galactosidase used for
genetic complementation

MCS = multiple cloning site containing the recognition sequences of the listed restriction enzymes

T3 and T7 = promoter sequences from phages T3 and T7.

Figure 2 shows the commercial plasmid vector pGEX-3x (Pharmacia Biotech) that was used for cloning of the PCR fragments corresponding to amino acid residues 27 to 158 of hMSH2 and 750 to 928 of GTBP (SEQ ID NO:1). Primers used for amplification were: ⁵ CGGGATCCCCCGGAGAAGCCGACCACCAC³ C and 5'CGGAATTCCTGGCCATCAACTGCGGACAT3' for codons 27 to 158 of 5'CGGAATTCTCAACTCGTATTCTTCTG3' 5'CGGGATCCCCCTTGAGAGGCTACTCAGT3' for codons 750 to 928 of GTBP. The PCR products, identified respectively as SEQ ID NO: 13 and 14 were cloned between the BamHI and EcoRI The expression products, in the form polypeptides fused with glutathione-S-transferase, were purified by affinity chromatography on a commercial glutathione matrix (Pharmacia Biotech) as directed by the manufacturer. The pure fusion proteins were used for the immunization of New Zealand White SPF female rabbits by standard protocols as reported in the publication Antibodies: A Laboratory Manual (1988) eds. Harlow and Lane, Cold Spring Harbor Laboratories Press.

Figure 3 shows an alignment of the amino acid sequences of the conserved C-terminal regions of the four mismatch binding proteins, i.e. GTBP (H. sapiens), hMSH2 (H. sapiens), MSH2 (S. cerevisiae) and MutS (E. coli). Identical residues are in black boxes, conserved ones in shaded boxes. Sequences reported in the alignment correspond to entries MSH2 YEAST (MSH2) and MUTS_ECOLI

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(MutS) in the SwissProt databank, or the coding region of GenBank entry HSU04045 (hMSH2). The alignements show that a high degree of conservation exists among the three homologs, with the C-terminal part of the protein being particularly highly conserved. GTBP can therefore be considered a new member of the MSH family.

Figure 4 shows the sequence homology, at the protein level, between pairs of MSH family members. Section a shows the matrix obtained from the alignment of GTBP (on the abscissa) with the yeast GTBP homolog (GenBank accession number Z47746, on the ordinate); proteins show comparable length and a significant homology is evident throughout their whole sequence. Section b shows the matrix obtained from the alignment of yeast MSH2 (on the ordinate) with GTBP (on the abscissa); the proteins show different lengths and most of the homology is confined to the C-rerminal regions of the two sequences. Section c shows the matrix obtained from the alignment of human MSH2 protein (on the ordinate) with GTBP (on the abscissa); the proteins show different lengths and, also in this case, most of the homology is confined to the C-rerminal regions of the two sequences. Section d shows the matrix obtained from the alignment of human hMSH2 protein (on the ordinate) with the yeast MSH2 (on the abscissa); the two proteins show comparable length and the homology is evident throughout the entire sequence.

Figure 5 shows the effect of selective anti-hMSH2 and anti-GTBP antisera on the formation of the specific mismatch-binding complex. Pre-incubation of HeLa nuclear extracts with either antiserum prior to addition of the G/T heteroduplex DNA probe results in a diminuition of the specific band in the gel-shift assay, an effect not observed when the respective pre-immune sera were used. This figure proves that both hMSH2 and GTBP are present in the mismatch-binding factor. This gel-shift analysis was carried out as described in ref.15, except that

nuclear extracts were used (25). The antisera were added to the reaction mixtures 20 min prior to the addition of the radioactively-labelled probe. The figure is an autoradiogram of a native 6% polyacrylamide gel run in Tris-acetate/EDTA (TAE) buffer prepared according to Maniatis et al., Molecular cloning: a laboratory manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y., 1982.

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Figure 6 shows that the mismatch-binding activity can be reconstituted using GTBP and hMSH2 obtained using an in vitro translation system. The procedure followed to generate in vitro transcripts of the hMSH2, Cl and FLY5 coding sequences was as follows: The DNA region encoding hMSH2 was inserted into pCite-1; Cl and FLY5 ORFs were introduced into pCite-2b (Novagen). In vitro transcription and translation reactions were carried out as described in ref. 26, including a mock translation 35_{S-labeled} added DNA. reaction in the absence of products were analysed translation on a polyacrylamide gel treated with Amplify (Amersham), dried and autoradiographed. Gel-shift assays were performed as described in ref. 15. Aliquots of 5 µl of the single in vitro translation reactions were tested; in the premixing experiments, 2.5 µl of each of the two translation reactions were mixed and incubated for 15 min at room temperature before the addition of the probe. AMP at a concentration of 5 mM was included in all the DNA binding reactions so as to overcome the effect of ATP in the reticulocyte lysates, which prevents the formation of mismatch-specific protein-DNA complexes, according ref. 16. Section a is an autoradiogram of a denaturing 7.5% SDS-polyacrylamide gel showing that translation of hMSH2, GTBP (Cl) and FLY5 mRNAs in a reticulocyte lysate system (Promega) gave rise to expected polypeptides of 113, 142 and 122 kDa, respectively. Section b shows the gel-shift analysis which demonstrates the binding of the in vitro-translated proteins to the G/T heteroduplex. The

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figure is an autoradiogram of a native 6% polyacrylamide gel run in TAE buffer.

Figure 7 shows that mismatch binding activity is absent from cell extracts lacking GTBP or hMSH2. The experiment is based on the analysis of two cell lines LoVo cells derived from CRC: contain a homozygous deletion of hMSH2 alleles and do not exhibit G/T binding activity (13), while neither hMSH2 allele is mutated in DLD1 cells, in spite of the fact that also this cell line lacks G/T binding activity. Section a shows a gel-shift assay showing that extracts of LoVo and DLDl fail to make mismatch-specific complexes. The G/C and G/T probes were obtained as described previously (15). Experimental The figure in Figure 6. conditions were as is autoradiogram of a native 6% polyacrylamide gel run in TAE buffer. Section b shows the Western blot analysis of extracts from Hela, LoVo and DLDl cells. The protein bands were visualized using an alkaline phosphataseconjugated anti-rabbit IgG system (Promega) as directed by the manufacturer. In the two left lanes, the anti-GTBP and anti-hMSH2 antisera were used alone with the HeLa extract to demonstrate their selectivity for the 160 and 100 kDa proteins, respectively. In the remaining lanes, both antisera were used together. Control HeLa cells revealed the presence of both hMSH2 and GTBP. contrast, the two CRC-derived tumor cell lines LoVo and DLD1 were completely devoid of full-length hMSH2 and GTBP, respectively. The amounts of hMSH2 in DLD1 cells and GTBP in LoVo cells were considerably lower than in HeLa cells. Since hMSH2 and GTBP bind heteroduplex DNA as a complex, the lack of one of the two proteins may cause instability of the second component of the complex.

Figure 8, part a, shows the experimental approach followed to discover the amino-terminal region of GTBP (from amino acid 1 to 68 of SEQ ID NO:15). Using the 5' RACE method(Rapid Amplification cDNA Ends, given in detail in the publication Nicolaides, N.C. et al.

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Genomics, 29: 229-234, 1995 and Nicolaides N.C. et al. Genomics, 30: 195-206, 1995) it is possible to determine the sequence upstream of the amino acid Ala in position 1 of SEQ ID NO:1. Initially, a pair of oligonucleotides was used that pairs with the sequence given in SEQ ID NO:12 from nucleotide 114 to 133 (primary oligonucleotide nucleotide 56 to 74 (secondary from The PCR reaction products were oligonucleotide B). sequenced and it was possible to determine that the capable of encoding amplification product was polypeptide DAAWSEAGPGPR, corresponding to amino acids 46-58 of the amino-terminal domain of GTBP as indicated in SEQ ID NO:15. Using a further two oligonucleotides, sequence was deduced from the initial RACE, complementary to the sequence given in SEQ ID NO:16 from nucleotide 188 to 204 (primary oligonucleotide C) and 169 185 (secondary oligonucleotide to oligonucleotide D) it was possible to amplify the GTBPcoding region 5' by-passing the methionine in position 1 of the amino acid sequence given in SEQ ID NO:15. The amplified clone, termed KMN, contained the entire nucleotidic sequence given in SEQ ID NO:16. analysis of leucocyte cDNA is shown in lanes 2 and 5, that of placenta cDNA in lanes 3 and 6. The products of lanes 1 to 3 derive from sequenced amplifications with oligonucleotides A and B, those in lanes 4 to 6 derive from sequenced amplifications with oligonucleotides C and Lanes 1 and 4 are the negative controls (absence of template). The molecular weight markers are indicated at the side.

shows expression of the figure 8 b of transcript encoding the protein GTBP using RT-PCR (PCR preceded by inverse transcription on RNA templates). RT-PCR was carried out using a synthetic oligonucleotide which paired with the sequence given in SEQ ID NO:12 from 114 to 133 in the inverse transcription nucleotide amplification reaction followed by with an

oligonucleotide with a sequence equal to the end 5' of the GTBP transcript, that is 5'GGTGCTTTTAGGAGCCCCG3'.

The RNA used as a mold template taken from HeLa cells (lane 2) placenta (lane 3) leucocytes (lane 4) and cells from the colon (lane 5); these were incubated with (+ symbol on the lane) or without (- symbol on the lane) inverse transcriptase and then made to underto PCR. Where no cDNA was produced, as the reverse transcription reaction did not occur, it was not seen to be amplified. Lane 1 is the negative control without RNA.

Detailed description

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In view of the potential and varied roles for mismatch binding proteins in the repair of information and their effects on disease state, such as tumor cell transformation and proliferation, metastases, and the paucity of understanding of the molecules and agents that selectively effect or modulate the activities of these proteins there exists a need in the art for compounds and agents with effector and modulator activity and methods to identify these and related compositions and agents. Further, such agents can serve as commercial research reagents for control of nucleic acid repair, and other GTBP-related conditions. Despite progress more defined model developing a of the molecular mechanisms underlying nucleic acid repair, significant methods applicable to assessing predisposition to cancer and or to its treatment have evolved. The hMSH2/GTBP heterodimer is necessary for the correction of base/base mispairs and one nucleotide loops. Genomic instability in tumor-derived cell-lines lacking GTBP demonstrates itself mainly in the form of small differences (e.g. in runs of A) rather than large changes in CA repeats, characteristic of phenotypes associated with the four known CRC loci hMSH2, hMLH1, hPMS1 and hPMS2. Cancers displaying mutator phenotypes with a low degree of microsatellite instability (20-24) may be associated with a malfunction of GTBP. It is a

discovery of the present invention that mutational events associated with tumorigenesis in CRC are due to defects in the GTBP gene.

Novel compositions comprising generic sequences encoding the GTBP protein, as well as fragments derived therefrom are provided, together with recombinant proteins produced using the genomic sequences and methods of using these compositions.

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Exemplary amino acid and DNA sequences invention are set forth in SEQ ID NO: 1 - SEQ ID NO:15 and in SEQ ID NO: 12 -SEQ ID NO: 16. abbreviations for nucleotides and amino acids are used in the Figures and elsewhere in this specification. GTBPparticularly preferred polypeptides are derived embodiments of the invention, although variations based on the specific sequences of these polypeptides are also part of the present invention. In its broader aspects, the invention (as it pertains to polypeptides per se) any polypeptide selected includes from the consisting of:

(i) any protein having an amino acid sequence which is at least 85% homologous to the amino acid sequences of SEQ ID NO: 1, SEQ ID NO:15 and the combination thereof, and, (ii) fragments thereof comprising at least 10 consecutive amino acids located within the amino acid sequences of SEQ ID NO: 1, SEQ ID NO:15 and the combination thereof, wherein the polypeptide is capable of binding to an antibody specific for GTBP.

In the genetic engineering aspects of the present invention, specific coding sequences as set forth in SEQ ID NO: 12, SEQ ID NO:16 and the combination thereof, which correspond to the preferred polypetides are themselves preferred.

Equivalent and complementary DNA and RNA sequences (see below for definitions of these terms) are likewise preferred. In its broader aspects, the genetic engineering aspects of the present invention include any

recombinant DNA or RNA molecule comprising a DNA sequence encoding GTBP itself or GTBP-derived protein according to SEQ ID NO: 1 or a corresponding DNA or RNA sequence, or a subsequence thereof comprising at least 10 nucleotides. The present invention also focuses on diagnostic methodologies aimed to detect loss of GTBP function in humans and consequent predisposition to neoplasia.

Defintion of terms

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A number of terms used in the art of genetic engineering and protein chemistry are used herein with the following defined meanings.

Two nucleic acid fragments are "homologous" if they hybridizing of to one another hybridization conditions described in Maniatis et al., (1982), Molecular cloning: a laboratory manual. Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y., pp. 320-325. By using the following wash conditions --2 xSSC, 0.1% SDS, room temperature twice, 30 minutes each; then 2 x SSC, 0.1% SDS, 50° C once, 30 minutes; then 2 x SSC, room temperature twice, 10 minutes each-- homologous sequences can be identified that contain at most about 25-30% base pair mismatches. More preferably, homologous nucleic acid strand contains 15-25% base pair mismatches, even more preferably 5-15% base pair mismatches. These degrees of homology can be selected by using more stringent wash conditions for identification of clones from gene libraries (or other sources of genetic material), as is well known in the art.

Two amino acid sequences are homologous if there is a partial or complete identity between their sequences. For example, 85% homology means that 85% of the amino acids are identical when the two sequences are aligned for maximum matching. Gaps (in either of the two sequences being matched) are allowed in maximizing matching gap lengths of 5 or less are preferred with 2 or less being more preferred.

Alternatively and preferably, two protein sequences (or polypeptide sequences derived from them of at least 30 amino acids in length) are homologous, as this term is used herein, if they have an alignment score of more than 5 (in standard deviation units) using the program ALIGN with the mutation data matrix and a gap penalty of 6 or greater (Dayhoff, M.O., in Atlas of Protein Sequence and Structure, 1972, volume 5, National Biomedical Research Foundation, pp. 101-110, and Supplement 2 to this volume, pp. 1-10). The two sequences or parts thereof are more preferably homologous if their amino acids are greater than or equal to 50% identical when optimally aligned using the ALIGN program.

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A DNA fragment is "derived from" a GTBP-encoding DNA sequence if it has the same or substantially the same base pair sequence as a region of the coding sequence for GTBP protein molecule.

"Substantially the same" means, when referring to biological activities, that the activities are of the same type although they may differ in degree. When referring to amino acid sequences, "substantially the same" means that the molecules in question have similar biological properties and preferably have at least 85 % homology in amino acid sequences. More preferably, the amino acid sequences are at least 90% identical. In other uses, "substantially the same" has its ordinary English language meaning.

A protein is "derived from" GTBP if it has the same or substantially the same amino acid sequence as a region of the GTBP protein molecule. By polypeptide derivatives of GTBP protein is meant polypeptides differing in length from the natural protein and containing five or more amino acids in the same primary order as found in the protein as obtained from a natural source. Polypetide molecules having substantially the same amino acid sequence as the natural protein but possessing minor amino acid substitutions which do not significantly

affect the ability of the protein or polypeptide to interact with protein-specific molecules, such as antibodies and nucleic acids are within the definition as derived from GTBP. Derivatives include glycosylated forms, aggregative conjugates with other protein molecules and covalent conjugates with unrelated chemical moieties. Covalent derivatives are prepared by linkage of functionalities to groups which are found in the amino acid chain or at the N-or C-terminal residue by means known in the art.

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GTBP-specific molecules include polypeptides such as specific for the antibodies that are protein polypeptide containing the naturally occurring GTBP amino acid sequence. By "specific binding polypetide" intended polypeptides that bind with GTBP protein and its derivatives and which have a measurably higher binding affinity for the target polypeptide than for other polypetides tested for binding. Higher affinity by a factor 10 is preferred, more preferably by a factor of 100. Binding affinity for antibodies refers to a single binding event (i.e., monovalent binding of an antibody molecule). Specific binding by antibodies also means that binding takes place at the normal binding site of the molecule's antibody (at the end of the arms in the variable region).

As discussed above, minor amino acid variations from the natural amino acid sequence of GTBP protein are contemplated; in particular, conservative amino acid replacements are contemplated. Conservative replacements are those that take place within a family of amino acids that are related in their side chains. Genetically encoded amino acids are generally divided into four families: (I) acidic = aspartate, glutamate; (2) basic = lysine, arginine, histidine; (3) non-polar = alanine, isoleucine, proline, phenylalanine, valine, leucine, methionine, tryptophan; and (4) uncharged polar, = glycine, asparagine, glutamine, cystine, serine,

Phenylanine, tryptophan, tyrosine. threonine, tyrosine are sometimes classified jointly as aromatic amino acids. For example, it is reasonable to expect that an isolated replacement of a leucine with an isoleucine or valine, an aspartate with a glutamate, a theonine with a serine, or a similar replacement of an amino acid with a structurally related amino acid will not have a major effect on the binding properties of the resulting molecule, especially if the replacement does not involve amino acid at a binding site involved in an interaction of GTBP or its derivatives with an antibody or with a specific DNA recognition sequence. Whether an amino acid change results in a functional peptide can readily be determined by assaying the specific binding properties of the polypeptide derivative.

Isolation of cDNA encoding GTBP protein

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Isolation of nucleotide sequences encoding GTBP protein involves creation of a cDNA library prepared from full-length mature messenger RNA extracted from cultured cells or tissues. Evidence is provided that GTBP is conserved over a broad evolutionary range, thus allowing the isolation of GTBP homologs from the genomes of phylogenetically distant species, i.e. from mammals to yeasts to bacteria.

Genetic libraries can be made in either eukaryotic prokaryotic host cells. Widely available cloning vectors such as plasmids, cosmids, phage, YACs and the like can be used to generate genomic libraries suitable for the isolation of nucleotide sequences encoding GTBP protein or portions thereof. Useful methods for screening genetic libraries for the presence of GTBP protein sequences include the preparation nucleotide oligonucleotide probes based on the sequence information provided in SEQ ID NO: 1 and SEQ ID NO: 15 (after decoding of the amino acid sequence) as well as in SEQ ID NO:12 and SEO ID NO: 16 (directly derived from the encoding DNA) of this patent. By employing the standard

triplet genetic code, oligonucleotide sequences of about 17 base pairs or longer can be prepared by conventional in vitro synthesis techniques. The resultant nucleic acid sequences can be subsequently labeled with radionuclides, enzymes, biotin, fluorescers or the like, and used as probes for screening the libraries.

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Additional methods of interest for isolating GTBP protein-encoding nucleic acid sequences include screening of genetic libraries for the expression of GTBP protein or fragments thereof by means of GTBP protein-specific antibodies, either polyclonal or monoclonal. Moreover, a selection method advisable for the screening of GTBP libraries cloned in conventional expression vectors is based on the specific binding of the protein polypeptides contained therein) to heteroduplex containing G/T molecules mimatches. A particularly preferred technique for isolating homolog proteins from related species or strains involves the use of degenerate primers based on partial amino acid sequences of GTBP protein and the polymerase chain reaction (PCR) amplify gene segments between the primers. A similar approach can also be applied to generate double stranded molecules after amplification of appropriate primers and polymerases. The gene can than be isolated using a specific hybridization probe based on the amplified gene segment, which is then analyzed for appropriate expression of the protein.

The nucleotide sequence of the isolated genetic material which encodes GTBP protein can be obtained by sequencing the non-vector nucleotide sequences of these recombinant molecules. Nucleotide sequence information can be obtained by employing widely used DNA sequencing protocols, such as Maxam and Gilbert sequencing, dideoxy nucleotide sequencing according to Sanger, and the like. Examples of suitable nucleotide sequencing protocols can be found in Berger and Kimmel, Methods in Enzymology Vol 52 Guide to Molecular Cloning Techniques, (1987) Academic

Press. Nucleotide sequence information from several recombinant DNA isolates, including isolates from both cDNA and genomic libraries, may be combined so as to provide the entire amino acid coding sequence of GTBP, as well as the nucleotide sequences of upstream and downstream nucleotide sequences.

Nucleotide sequences obtained from sequencing GTBP protein-specific genomic library isolates subjected to further analysis in order to identify regions of interest in the GTBP gene. These regions of include additional interest open reading promoter sequences, termination sequences, and the like. Analysis of nucleotide sequence information is preferably performed by computer. Software suitable for analyzing for regions of interest nucleotide sequences commercially available and includes, for example, DNASIS (Pharmacia Biotech). It is also of interest to use amino acid sequence information obtained from the sequencing of purified GTBP protein when analyzing new GTBP nucleotide sequence information so as to improve the accuracy of the nucleotide sequence analysis.

Expression of GTBP

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Isolated nucleotide sequences encoding GTBP protein can be used to produce purified GTBP protein or fragments thereof by either recombinant DNA methodology or by in vitro polypeptide synthesis techniques. By "purified" and "isolated" is meant, when referring to a polypeptide or nucleotide sequence, that the indicated molecule is present in the substantial absence of other biological macromolecules of the same type. The term "purified" as used herein preferably means at least 95% by weight, more preferably at least 99% by weight, and most preferably at least 99.8% by weight, of biological macromolecules of the same type present (but water, buffers, and other small molecules, especially molecules having a molecular weight of less than 1000, can be present).

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A significant advantage of producing GBTB protein by recombinant DNA techniques rather than by isolating from natural sources of GTBP protein is that equivalent quantities of GTBP protein can be produced by using less starting material than would be required for isolating GTBP protein from a natural source. Producing protein by recombinant techniques also permits protein to be isolated in the absence of some molecules normally present in cells that naturally produce GTBP protein. It is also apparent that recombinant techniques can be used to produce GTBP protein polypeptide derivatives that are not found in nature, such as the variations described above.

GTBP protein and polypeptide derivatives of GTBP protein can be expressed by recombinant techniques when a DNA sequence encoding the relevant molecule functionally inserted into a vector. By "functionally inserted" is meant in proper reading frame orientation, as is well understood by those skilled in the art. Typically, the GTBP protein gene will inserted downstream from a promoter and will be followed by a stop codon, although production as a hybrid protein followed by cleavage may be used, if desired. In general, host-cell-specific sequences improving the production yield of GTBP protein and GTBP polypeptide derivatives will be used, and appropriate control sequences will be added to the expression vector, such as enhancer polyadenylation sequences, sequences, and ribosome binding sites.

Two basic types of expression are contemplated: (i) expression in mammalian cells so as to overcome a deficiency in an individual having insufficient GTBP, and (ii) expression for the purpose of providing GTBP for purpose irrelevant to the host in which expression occurs, such as production of diagnostic tests for GTBP deficiency.

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Production of genetic constructs for transformation of human cells

With the goal of expression in human cells, a gene construct will be prepared and used to transform human cells. Several strategies and vectors have been developed for the expression of proteins in animal cells. For example BK-SV40 hybrid vectors have been constructed . These vectors can be maintained in cultured human cells double-stranded DNA multicopy extrachromosomal replicons. One exemplary vector consists of the SV40 promoter controlling the expression of resistance gene (the selectable marker) and the MMTV promoter regulated by the DRE enhancer sequence which controls the expression of the cloned gene. In any case, the foreign construct will usually translational initiation transcriptional and and termination signals, with the initiation signals 5' to the gene and termination signals 3' to the gene of interest, altough linear DNA can be delivered to a host where recombination occurs for insertion into the host genome. Expression under the control of the native promoter can thus be achieved by replacing the defective gene with the linear DNA encoding GTBP by making use of cellular processes, e.g. homologous recombination. transcriptional initiation region which includes the RNA polymerase binding site (promoter) may be native to the host or may be derived from an alternative source, where the region is functional in the host. The transcriptional initiation regions may not only include polymerase binding site, but also regions providing for the regulation of the transcription. The 3' termination region may be derived from the same gene trancriptional initiation region or a different gene. For example, where the gene of interest has a trascriptional termination region functional in the host species, that region may be retained within the gene.

An expression cassette can be constructed which will include transcriptional initiation region, the GTBP protein gene under the transcriptional control of the trascription initiation region, the initiation codon, the coding sequence of the gene, with or without introns, and translational stop codons, followed the transcriptional termination region, which will include the terminator, and may include a polyadenylation signal sequence, and other sequences associated transcriptional termination. The direction is 5' to 3' same as the direction of transcripition. The cassette will usually be less than about 10 kb, frequently less than about 6 kb, usually being at least about 5 kb.

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When the expression product of the gene is to be located other than in the cytoplasm, the gene will usually be constructed to include particular amino acid sequences which result in translocation of the product to a particular site, which may be an organelle, such as the nucleus, or may be secreted into the external environment of the cell. Various secretory leaders, membrane integrator sequences, and translocation sequences for directing the peptide expression product to a particular site are described in the literature.

One or more cassettes may be involved, where the cassettes may be employed in tandem for the expression of independent genes which may express products independently of each other or may be regulated concurrently, where the products may act independently or in conjunction, e.g. GTBP and hMSH2.

The expression cassette will normally be carried on a vector having at least one replication system. For convenience, it is common to have a replication system functional in *E. coli* such as ColE1, pSC101, pACYC184, or the like. In this manner, at each stage after each manipulation, the resulting construct may be cloned, sequenced, and the correctness of the manipulation determined. In addition, or in place of the *E. coli*

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replication system, a broad host range replication system may be employed, such as the replication systems of the Pl incompatibility plasmids, e.g. RK2, RP1, RP4 and R68.

In addition to the replication system, there will frequently be at least one marker present, which may be uselful in one or more hosts, or different markers for individual hosts. That is, one marker may be employed for selection in a prokaryotic host, while another marker may be employed for selection in a eukaryotic host. Various genes which may be employed include neo (neomycin-kanamycin resistance), choramphenicol acetyltransferase (cat), b lactamase (bla), b galactosidase etc.

The various fragments comprising the various constructs, expression cassettes, markers, and the like may be introduced consecutively by restriction enzyme cleavage of an appropriate replication system, and insertion of the particular construct or fragment into the available size. After ligation and cloning the vector may be isolated for further manipulation. All of these techniques are amply exemplified in the literature and find particular exemplification in Maniatis et al., Molecular cloning: a laboratory manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y., 1982.

Transformation of mammalian cells and gene therapy

Once the vector is completed, the vector may be mammalian cells. Techniques introduced into transfection. transforming mammalian cells include microinjection, liposome-based delivery Transfection of cultured human cells is the most commonly used method and can be achieved by standard protocols which involve either incubation of cells with DNA that has been co-precipitated with calcium phosphate or DEAEdextran or electroporation with purified transfecting DNA. In other systems, a genetically modified virus, a liposome or a microinjection can also be used to deliver foreign DNA to human recipient cells. Once the GTBP gene has been introduced into the defective cell, it can 5

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complement the genetic defect, restoring the normal phenotype. This methodology, when used to remediate genetic defects in individuals, goes under the name of gene therapy. At least two strategies for implementing somatic cell gene therapy have emerged and could be applied to correct GTBP genetic defects: ex vivo and in vivo gene therapy. Usually, the ex vivo gene therapy involves the following procedures:

- collect the cells from an affected individual
- correct the genetic defect by gene transfer
- select and grow the genetically corrected (remedial) cells
- infuse or transplant corrected cells back into the patient.

Vectors derived from retroviruses are often used to stably maintain and persintently express the remedial gene in the corrected cell.

In vivo gene therapy entails the direct delivery of remedial gene into the cell of a particular tissue of a prospective patient. The wild-type protein can be cloned into various benign viruses and delivered to target defective cells in an in vivo infection. Vectors derived from adenovirus, herpes simplex virus and certain retroviruses are excellent candidates for in vivo gene therapy. Methods and prospectives of gene therapy have been reviewed by Mulligan (1993), Science 260:926-932.

Diagnostic methods using antigens

Typically, methods for detecting analytes such as binding proteins of the invention are based on immunoassays. Immunoassays can be conducted to determine the presence or absence of GTBP in host cells. Such techniques are well known and need not be described here in detail. Examples include both heterogeneous and homogeneous immunoassay techniques. Both techniques are based on the formation of an immunological complex between the binding protein and a corresponding specific antibody. Heterogeneous assays for GTBP typically use a

specific monoclonal or polyclonal antibody bound to solid surface, e.g. in sandwich assays. Homogeneous assays, which are carried out in solution without the presence of a solid phase, can be used, for example, by determining the difference in enzyme activity brought on by binding of free antibody to an enzyme-antigen conjugate. A number of suitable assays are disclosed in U.S. Patent Nos. 3,817,837, 4,006,360, and 3,996.34545.

The solid surface reagent in the above assay prepared by known techniques for attaching protein material to solid support material, such as polymeric beads, dip sticks, or filter material. These attachment methods generally include non-specific adsorption of the protein to the support or covalent attachment of the protein, typically through a free amine group, to a chemically reactive group on the solid support, such as an activated carboxyl, hydroxyl, or aldehyde group.

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In a second diagnostic configuration, known as a antibody binding to assay, an homogeneous produces some change in the reaction medium which can be directly detected in the medium. Known general types of homogeneous assays proposed heretofore include (a) spinlabeled reporters, where antibody binding to the antigen is detected by a change in reported mobility (broadening of the spin splitting peaks), (b) fluorescent reporters, where binding is detected by a change in fluorescence emission, (c) enzyme reporters, where antibody binding effects enzyme/substrate interactions, and (d) liposomebound reporters, where binding leads to liposome lysis and release of encapsulated reporter. The adaptation of these methods to the protein antigen of the present invention follows conventional methods for preparing homogeneous assay reagent.

In each of the assays described above, the assay method involves reacting the tissue extract from a test individual with an antibody and examining the sample for the presence of bound antigen. The examination may

involve attaching a labelled anti-GTBP antibody to the primary complex formed between GTBP and the immobilized antibody and measuring the amount of reporter bound to the solid support, as in the first method, or may involve observing the effect of antibody binding on a homogeneous assay reagent, as in the second method.

Production of specific binding proteins

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GTBP, in its native or chemically modified form, or polypeptide derivatives thereof, or specific complexes with other polypeptides may be used for producing antibodies, either monoclonal or polyclonal, specific to GTBP or polypeptide derivatives thereof, or to GTBP complexes with other polypeptides. Antibodies specific are protein produced by immunizing appropriate vertebrate host, e.g., rabbit or mouse, with purified GTBP protein or polypeptide derivatives of GTBP by themselves or in conjunction conventional adjuvant. Usually, two or more immunizations will be involved, and blood or spleen will be harvested a days after the last injection. For few polyclonal the immunoglobulins be precipitated, antisera, can purified of isolated and by a variety standard techniques, including affinity purification using GTBP protein attached to a solid surface, such as a gel or beads in an affinity column. For monoclonal antibodies, splenocytes will normally be fused with immortalized lymphocyte, e.g., a myeloid cell line, under selective conditions for hybridoma formation. hybridomas can then be cloned under limiting dilution conditions and their supernatants screened for antibodies having the desired specificity. Techniques for producing antibodies are well known in the literature and are exemplified by the publication Antibodies: A Laboratory Manual (1988) eds. Harlow and Lane, Cold Spring Harbor Laboratories Press, and U.S. Patent Nos. 4,381,292, 4,451,570, and 4,618,577.

GTBP diagnostic application using genetic probes

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The genetic material of the invention can itself be used in numerous assays as probes for genetic material present in an individual. The analyte can be a nucleotide sequence which hybridizes with a probe comprising a sequence of at least about 16 consecutive nucleotides, usually 30 to 200 nucleotides, up to substantially the full sequence of the gene as shown in SEQ ID NO: 12. The analyte can be RNA or DNA. The sample is typically a DNA or an RNA molecule extracted by the patient's tissue. In order to detect an analyte, where the analyte hybridizes to a probe, the probe may contain a detectable label. Particularly preferred for use as a probe are sequences up to about 3200 consecutive nucleotides (for example from nucleotide 1 to nucleotide 3000 of SEQ ID NO: 12 and from nucleotide 1 to nucleotide 204 of SEQ ID NO:16) since these sequences appear to be particularly specific for GTBP.

One method for amplification of target acids, for later analysis by hybridization assays, known as the polymerase chain reaction or PCR technique. The PCR technique can be applied to detect sequences of the invention in suspected samples using oligonucleotide primers spaced apart from each other and based on the genetic sequence set forth in SEQ ID NO: 12 and SEQ ID NO:16. The primers are complementary to opposite strands a double-stranded DNA molecule and are typically separated by from about 50 to 450 nt or more (usually not more than 2000 nt). This method entails preparing the specific oligonucleotide primers and then repeated cycles of target DNA denaturation, primer binding, and extension with a DNA polymerase to obtain DNA fragments of the expected length based on the primer spacing. Extension products generated from one primer serve as additional target sequences for the other primer. The degree of amplification of a target sequence is controlled by the number of cycles that are performed and is theoretically calculated by the simple formula 2ⁿ where n is the number

of cycles. Given that the average efficiency per cycle ranges from about 65% to 85%, 25 cycles produce from 0.3 to 4.8 million copies of the target sequence. The PCR method is described in a number of publications, including Saiki et al., Science (1985) 230:1350-1354; Saiki et al., Nature (1986) 324:163-166; and Scharf et al., Science (1986)233:1076-1078. Also see U.S. Patent Nos. 4,683,194; 4,683,195; and 4,683,202.

The invention includes a specific diagnostic method for determination of GTBP, based on selective amplification of GTBP-protein-encoding DNA fragments. This method employs a pair of single-stranded primers derived from non-homologous regions of opposite strands DNA duplex fragment having a sequence described by combining the sequences SEQ ID NO: 16 and SEQ ID NO:12. These "primer fragments" represent one aspect of the invention. The method follows the process amplifying selected nucleic acid sequences disclosed in U.S. Patent No. 4,683,202, as discussed above.

Mutations in the *GTBP* gene can be detected by restriction enzyme analysis of the amplification product or by direct sequencing. Also, alterations in *GTBP* sequence can be revealed by Southern hybridization with probes encompassing part or the entire sequences of SEQ ID NO: 12 and SEQ ID NO:16.

Single-stranded DNA probes complementary to the wild-type GTBP-coding sequence can also be hybridized to RNA extracted from tissues or cells of human patients and used to detect mutations in the mature GTBP gene transcript by enzymatic digestion of heteroduplexes at the level of mismatches. These and other techniques aimed to identify variations in gene sequences from wild-type GTBP are extensively reported in the literature and well established in the scientific community.

Binding assays involving GTBP

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The presence of an altered GTBP protein can be detected by the use of binding assays based on the specific recognition of G/Tmismatches by GTBP. synthetic double-stranded 34-mer oligonucleotide G/T mispair is containing prepared and labelled substantially as reported (15). Cell extracts can be prepared as reported in current literature (e.g. ref 25 and refs. therein). The cell extract (1-10 micrograms of nuclear proteins) can be incubated with the heteroduplex oligonucleotide at room temperature for 30 minutes to allow GTBP binding to the G/T mismatch. The mixture can then be loaded on a gel prepared as reported in Figure 6. Alterations in GTBP mass or affinity for the substrate can be evidenced by an altered electrophoretic mobility.

15 <u>Deposits</u>

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Strains of E. coli TOP10 - transformed using the plasmids pBluescript SK-/C1 and pCite-2b/C1 coding respectively for the protein GTBP from the amino acid 1 to the amino 1292 SEQ NO:1 and using the plasmid of ID pBluescript SK /FLYS coding for a GTBP protein from the amino acid 116 to the amino acid 1292 of SEQ ID NO:1 on 19/5/1995 with the National have been deposited Industrial and Marine Bacteria Collections of (NCIMB), Aberdeen, Scotland, UK, with accession numbers NCIMB 40742, NCIMB 40471 and NCIMB 40740 respectively. Moreover, a strain of E.coli TOP10 - transformed using the plasmid pBluescript SK /GTBP coding for the whole amino acid sequence of GTBP from the amino acid 1 to the amino acid 1360 (SEQ ID NO: 15 and SEQ ID NO:1) - has been 28/5/96 with the above deposited on institution with accession number NCIMB 40805.

Examples

As mentioned above, the inventors identified a mismatch-binding factor in HeLa cells (15), GTBP, which was shown to bind preferentially to heteroduplexes containing G/T mispairs. Purification of this DNA binding activity by G/T mismatch affinity chromatography yielded

a mixture of two proteins of apparent molecular weights of 100 and 160 kDa (16), which indicates that the mismatch-specific complex is composed of two proteins. The 100 kDa constituent of the complex is hMSH2 (17) while the second component is GTBP. Examples regarding the identity and function of GTBP are reported below.

Example 1

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The present example shows that the GTBP protein sequence, as reported by combining the sequences SEQ ID NO:15 and SEQ ID NO: 1, contains seven subsequences which correspond to polypeptides obtained after proteolytic cleavage of the 160 kDa DNA-binding protein termed GTBP. These subsequences are indicated as SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7 and SEQ ID NO: 8. The 160 kDa protein was purified as reported in ref. 16. The fractions containing the G/Tspecific mismatch binding activity were loaded onto a preparative SDS-PAGE gel and the 100 and 160 kDa bands were excised following staining with Coomassie Blue. The proteins were digested in the gel matrix either with trypsin (100 kDa protein, Promega Corporation, UK), or with Achromombacter lyticus endopeptidase lys-C (160 kDa protein, Wako Chemicals GmbH, Germany). The proteolytic peptides were recovered by sequential extractions and separated by tandem hplc on a Hewlett-Packard 1090M with diode array detection. Anion-exchange and phase columns were connected in series, essentially as described by H. Kawasaki and K. Suzuki, Anal. Biochem. 186, 264 (1990). Fractions were collected and applied directly to an Applied Biosystems 477A pulsed-liquid automated sequencer modified as described by N.F. Totty, M.D. Waterfield and J.J. Hsuan, Protein 1. 1215 (1992). Microsequencing yielded seven proteolytic peptides whose sequences have been designated as SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7 and SEQ ID NO: 8. Example 1B

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The present example shows that the protein GTBP contains an amino-terminal domain corresponding to SEO ID NO:15. This region can be determined by analysis of the coding nucleotide sequence. The amino-terminal domain is integral part of the peptide GTBP itself, therefore the GTBP sequence must be understood to be the sequenced combination of SEQ ID NO:15 and SEQ ID: NO:1 with a total extension of 1360 amino acids. figure 8 shows the experimental approach followed to discover the amino-terminal region of GTBP (from amino acid 1 to 68 of SEQ ID NO:15). Using the 5' RACE method(Rapid Amplification cDNA Ends, given in detail in the publication Nicolaides, N.C. et al. Genomics, 229-234, 1995 and Nicolaides N.C. et al. Genomics, 30: 195-206, 1995) it is possible to determine the sequence upstream of the amino acid Ala in position 1 of SEQ ID Initially, a pair of oligonucleotides was used that pairs with the sequence given in SEQ ID NO:12 from nucleotide 114 to 133 (primary oligonucleotide A) from nucleotide 56 to 74 (secondary oligonucleotide B). The PCR reaction products were sequenced and it possible to determine that the amplification product was capable of encoding the polypeptide DAAWSEAGPGPR, corresponding to amino acids 46-58 of the amino-terminal domain of GTBP as indicated in SEQ ID NO:15. Using a further two oligonucleotides, whose sequence was deduced from the initial RACE, complementary to the sequence given in SEQ ID NO:16 from nucleotide 188 to 204 (primary oligonucleotide C) and from oligonucleotide 169 to 185 (secondary oligonucleotide D) it was possible to amplify the GTBP-coding region 5' by-passing the methionine in position 1 of the amino acid sequence given in SEQ ID The amplified clone, termed KMN, contained the entire nucleotidic sequence given in SEQ ID NO:16. analysis of leucocyte cDNA is shown in lanes 2 and 5, that of placenta cDNA in lanes 3 and 6. The products of lanes 1 to 3 derive from sequenced amplifications with

oligonucleotides A and B, those in lanes 4 to 6 derive from sequenced amplifications with oligonucleotides C and D. Lanes 1 and 4 are the negative controls (absence of template). The molecular weight markers are indicated at the side.

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Part b of figure 8 shows expression the transcript encoding the protein GTBP using RT-PCR (PCR preceded by inverse transcription on RNA templates). RT-PCR was carried out using a synthetic oligonucleotide which paired with the sequence given in SEQ ID NO:12 from nucleotide 114 to 133 in the inverse transcription followed by amplification with oligonucleotide with a sequence equal to the end 5' of the GTBP transcript, that is 5'GGTGCTTTTAGGAGCCCCG3'.

The RNA used as a mold was taken from HeLa cells (lane 2) placenta (lane 3) leucocytes (lane 4) and cells from the colon (lane 5); these were incubated with (+ symbol on the lane) or without (- symbol on the lane) inverse transcriptase and then made to underto PCR. Where no cDNA was produced, as the reverse transcription reaction did not occur, it was not seen to be amplified. Lane 1 is the negative control without RNA.

Example 2

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The present example shows that DNA regions internal to GTBP gene can be obtained by amplification with primers designed on the basis of the sequence of peptides deriving from proteolytic cleavage of the 160 kDa G/T-binding factor (SEQ ID NO: 2 to 8). Following the strategy of Lingner et al. (18) the inventors identified a unique DNA sequence encoding the central 8 amino acids of the peptide of SEQ ID NO: 6. Two degenerate primers corresponding to the N- and C-terminal amino acid sequences of the oligopeptide of SEQ ID NO: 6, i.e. the DNA sequences 5'GCGAATTCTAYGGNTTYAAYGC3' (SEQ ID NO: 9) and

 $^5\,^{\prime}\text{GCGGATCCTAYTGDATNACYTC}^3\,^{\prime}$ (SEQ ID NO: 10), where N=any nucleotide, Y=C or T and D=A, G or T

were used for PCR amplification on poly-A+ HeLa mRNA as described (18) except that the MgCl₂ concentration was 5 mM. The expected 67 bp fragment was eluted from an acrylamide gel, cloned into pBluescript SK- and sequenced (see. comments to SEQ ID NO: 9 and 10 for details). Two clones contained the correct sequence, corresponding to SEQ ID NO: 11, encoding the starting target peptide SEQ ID NO: 6..

Example 3

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The present example shows that DNA regions internal to GTBP gene can be identified by hybridization with a DNA probe designed on the basis of the sequence of peptides obtained upon proteolytic cleavage of the 160 kDa G/T-binding factor. The DNA sequence reported as SEQ ID NO: 11 was was labeled with ³²P by a standard kinase reaction (with T4 PNK and [q-32P]ATP as described by Maniatis et al., Molecular cloning: a laboratory manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y., 1982) in order to generate a double-stranded DNA probe. The labelled probe of SEQ ID NO: 11 was then used in the screening of a commercial oligo dT-primed cDNA library in phage lambda (HeLa S3 Uni-ZAP XR, Stratagene). positive clones were selected for further analysis. Clone C1 contained an insert of 3980 bp corresponding to SEQ ID NO: 12, with a continuous open reading frame from amino acid residue 1 to 1292 encoding a polypeptide of 1292 amino acids (SEQ ID NO: 1) and a calculated molecular mass of 142 kDa; clone FLY 5 contained sequences coding from aa residue 116 to 1292 (see comments to SEQ ID NO: 1 and 12).

As all seven peptides obtained from the microsequencing of the 160 kDa protein (SEQ ID NO: 2 to 8) could be found in SEQ ID NO: 1, it can be concluded that clone C1 encodes GTBP.

35 Example 4

The present examples shows that GTBP protein can be used as an antigen to produce highly specific antibodies

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which recognize GTBP but not hMSH2. PCR fragments corresponding to amino acid residues 27 to 158 of hMSH2 (SEQ ID NO: 13) and 750 to 928 of GTBP (SEQ ID NO: 14) were subcloned into the E. coli expression vector pGEX-3X (Pharmacia/LKB) and the recombinant proteins, in the form of fusion polypeptides with glutathione S-transferase, induced and isolated as recommended manufacturer, except that the final concentration of IPTG was 0.25 mM and induced cultures were harvested after 6 hours at 20°C. The fusion proteins were used immunization of New Zealand White S.P.F. female rabbits using standard protocols. (Charles River Co.) Two polyclonal antisera specifically immunoreactive to GTBP and hMSH2, respectively, were obtained and assayed as reported in Antibodies: A Laboratory Manual (1988) eds. Harlow and Lane, Cold Spring Harbor Laboratories Press (see Figures 2 and 5 for more details).

Example 5

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The following example shows that GTBP belongs to a class of DNA-repair proteins conserved over a wide 20 evolutionary range. Figure 3 shows the alignment of the amino acid sequences of the conserved C-terminal regions of the mismatch binding proteins GTBP (H. sapiens), hMSH2 (H. sapiens), MSH2 (S. cerevisiae) and MutS (E. coli). Identical residues are in black boxes, conserved ones in 25 shaded boxes. Sequences reported in the alignment correspond to entries MSH2 YEAST (MSH2) and MUTS ECOLI (MutS) in the SwissProt databank, or the coding region of GenBank entry HSU04045 (hMSH2). The alignment was carried 30 out using the GCG Pileup option. The figure was generated using Prettyplot. The alignment reveals a high degree of conservation at the C-terminal domain among all proteins. GTBP can thus be considered a new member of the Muts Homolog (MSH) family.

However, GTBP must be considered structurally distinct from MSH proteins, since the N-terminal domain (up to approximatively 1000 amino acids) of GTBP exhibits

remarkable divergency from MSH (human, yeast bacterial). This is particularly evident when the homology matrixes of hMSH2 versus MSH2 (Figure 4 section d) and GTBP versus hMSH2 (Figure 4 section c) or GTBP versus MSH2 (Figure 4 section b) are compared to one another. In contrast, clear evidence is provided that GTBP is conserved over a wide evolutionary range and that structural homologs of GTBP through the whole sequence can also be found , e.g. in yeast (GenBank accession number Z47746, Figure 4 section a).

Example 6

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The following example demonstrates that selective antisera recognize hMSH2 and GTBP bound to mismatched DNA in a complex. Figure 5 shows the effect of anti-hMSH2 and anti-GTBP antisera on the formation of the specific mismatch-binding complex. This gel-shift analysis was carried out as described (15), except that nuclear extracts were used (25). The antisera were added to the reaction mixtures 20 min prior to the radioactivelylabelled probe. The figure is an autoradiogram of a native 6% polyacrylamide gel run in TAE buffer. Preincubation of the HeLa nuclear extracts with either antiserum prior to the addition of the G/T heteroduplex probe resulted in the diminuition of the specific band in a gel-shift assay, an effect not observed when the sera were used. respective pre-immune This indicates that both proteins are present in the mismatchspecific factor. This finding also implies that extracts from cells lacking either protein are devoid of mismatchbinding activity.

Example 7

The following example shows that GTBP and hMSH2 can be expressed separately in a cell-free translation system. The inventors employed a hMSH2 cDNA clone (17) and the GTBP clones C1 and FLY5 as set forth in SEQ ID NO: 12. The C1 and FLY5 ORFs were introduced into pCite-2b. The hMSH2 ORF was inserted into pCite-1 (Novagen). In

and translation reactions vitro transcription carried out as described previously (26) including a mock translation reaction in the absence of added DNA. labeled translation products were analyzed on a SDSpolyacrylamide gel treated with Amplify (Amersham), dried and autoradiographed. The experiment was carried out using conditions recommended by the manufacturer. figure is an autoradiogram of a denaturing 7.5% SDSpolyacrylamide gel. As shown in Fig. 6 section a. translation of hMSH2, GTBP (C1) and FLY5 mRNAs in a reticulocyte lysate system (Promega) gave polypeptides of 113, 142 and 122 kDa respectively. Thus, translation of all three mRNAs gave rise to protein products of the expected size.

Example 8

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The following examples shows that GTBP binds G/T mismatches when complexed to hMSH2. This was achieved by testing the two polypeptides expressed in a cell-free translation system for their ability to bind mismatchcontaining substrates. Reconstitution of the mismatchbinding activity using in vitro translated GTBP and hMSH2 is shown in Figure 6 section b. The figure shows a gelshift analysis showing the binding of the in vitrotranslated proteins to the G/T heteroduplex. When GTBP and hMSH2 proteins were tested for mismatch binding activity, it was noted that expression of either protein alone has no effect on the intensity of the endogenous G/T-specific band present in the lysates at low levels. In contrast, mixing of the hMSH2 and GTBP translation products resulted in a reproducible increase in the intensity of the mismatch-specific band. This result is confirmed by using the GTBP cDNA clone FLY5, which encodes a truncated GTBP protein (see SEQ ID NO: 1 and 12). Mixing of hMSH2 and FLY5 translation products with the G/T probe gave rise to a new band with a faster electrophoretic mobility than the endogenous complex, such as would be expected of a smaller species. This

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Example 9

experiment provides convincing evidence that the human mismatch binding complex is composed of hMSH2 and GTBP.

Gel-shift assays were performed as described in (15). 5ml aliquots of the single in vitro translation reactions were tested; in the pre-mixing experiments, 2.5 ml of each of the two translation reactions were mixed and incubated for 15 min at room temperature before the addition of the probe. 5 mM AMP was included in all the DNA binding reactions so as to overcome the effect of ATP in the reticulocyte lysates, which prevents the formation of mismatch-specific protein/DNA complexes (16). The figure is an autoradiogram of a native 6% polyacrylamide gel run in TAE buffer.

Genetic alterations in mismatch repair genes such as hMSH2, hMLH1, hPMS1 and hPMS2 (1) are known to cause the hypermutability found in many forms of hereditary colorectal cancers (CRC). Here we report examples showing that different cell lines from CRC, which display contain mutated GTBP hypermutable phenotype, alleles which are expressed into non functional proteins. We also show that the spectrum of mutations found in these cell lines is different from that caused by the inactivation of hMSH2 or of other mismatch repair genes. The following examples confirm the role of GTBP in the maintenance of human genome integrity in vivo and provide an explanation for the mutator phenotype observed in different CRC.

The following example shows that mismatch binding activity is absent from extracts of LoVo and DLD1 cells, both derived from human CRC. LoVo cells contain a homozygous deletion in both hMSH2 alleles (13) while neither hMSH2 allele appears to be mutated in the cell line DLD1 (19). Extracts of LoVo and DLD1 cells fail to make mismatch-specific complexes as revealed by gel-shift assay shown in Figure 7 section a (probes were prepared as described previously (15) and experimental conditions were as in Figure 5). The figure is an autoradiogram of a

native 6% polyacrylamide gel run in TAE buffer showing the absence of specific DNA-protein complexes of expected molecular mass in both LoVo and DLD1 extracts. Based on this it appears evident that the DLD1 cell line must be devoid of GTBP. Confirmatory results were also obtained by direct screening of LoVo and DLD1 cell extracts with specific antibodies directed against GTBP and hMSH2. As expected, western blot analysis of HeLa extracts revealed the presence of equivalent amounts of hMSH2 and GTBP. In contrast, LoVo cells could be shown to lack hMSH2, and DLD1 extracts were completely devoid of full-length GTBP (Figure 7 section b). Interestingly, the amounts of hMSH2 in DLD1 and of GTBP in LoVo extracts were considerably lower than in the HeLa extracts. Our explanation for this finding is that hMSH2 and GTBP are unstable when not in a complex (16).

Example 10

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The CRC-derived cell line HCT15 contains a full length hMSH2 protein but shows hypermutable phenotype (19). To determine whether HCT15 had a mutation in the GTBP coding sequence, the RNA of this cell line was reverse transcribed with random hexamers and reverse transcriptase according to standard protocols (e.g., see Powell et al., New Engl. J. Med. 329, 1982, 1993). The cDNA was then amplified with PCR using primers specific for the GTBP-coding sequence. The oligonucleotides used 5'-PGAGGGTTACCCCTGG-3' primer ACACTGTAAGTCTGTGTACC-3' for codons 32 to 458, primers 5'-PAGTGAAGGCCTGAACAGCC-3' and 5'-AAGTCCAGTCTTTCGAGCC-3' for codons 219 to 858, and primers 5'-PGAGAGGGTTGATACTTGCC-3' and 5'-AGAAGTCAACTCAAAGCTTCC-3' for codons 692 to 1292 (where P denotes a T7 promoter sequence and a ribosomebinding site for translation initiation (26) and codon numbers are those reported in SEQ ID NO: 1 and SEQ ID NO: 12). To detect mutations in the GTBP-coding sequence, the amplification first transcribed products were and translated in vitro using a commercial kit (Promega).

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Analysis of translation products in a PAGE-SDS gel GTBP polypeptides from revealed truncated products, corresponding to regions located at codons 32-458 (5'-end of the gene) and 692-1292 (3'-end of the Sequencing of these PCR products using gene). system (SequiTherm Polymerase, commercial Epicentre revealed that truncations were Technologies) frameshift mutations. The deletion of nucleotide 664 (a C) at codon 222 changed a leucine to a termination codon and a substitution of nucleotides 3307-3312 (GATAGA) with (see SEQ ID NO: 12) created a new termination codon several bp downstream.

Example 11

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MT1 is an alkylation-resistant lymphoblastoid cell line with a biochemical deficiency similar to that of HCT15 (see Goldmacher et al., J. Biol. Chem., 261, 12462, 1986; Kat et al. Proc. Natl. Acad Sci USA, 90, 6424, 1993). To ascertain whether MT1 had a GTBP mutation, the RNA of this cell line was reverse transcribed with random hexamers and reverse transcriptase and the cDNA was then amplified with PCR using primers specific for the GTBPcoding sequence as reported above. In vitro transcription and translation of GTBP-coding sequence from MT1 did not reveal truncated GTBP polypeptide after electrophoretic of GTBP was therefore analysis. The coding region sequenced and two missense mutation were found in the GTBP cDNA. The first was an GAT to GTT transversion at codon 1145 of SEQ ID NO: 1, resulting in a substitution of aspartic acid with valine. The aspartic acid at codon 1145 is located in the putative DNA-binding domain of GTBP, and the identical amino acid is found at homologous positions in GTBP (H. sapiens), hMSH2 (H. sapiens), MSH2 (S. cerevisiae) and MutS (E. coli). This highly conserved amino acid residue is therefore necessary for activity and non conservative substitutions at residue cause dramatic refuction of GTBP funcionality. The second was a GTT to ATT transition, resulting in a

substitution of isoleucine to valine at codon 1193 of SEQ ID NO: 1.

The amplification products were cloned in the vector BLUESCRIPT SK⁻ and individual clones were sequenced using conventional protocols (Sequenase, USB). The two mutations were not found to be associated in a single clone, deriving thus from separate alleles.

Example 12

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A tumor cell line, termed 543X (from the patient's designation) derived was from CRC and displays hypermutable phenotype and microsatellite instability but no mutation in mismatch repair genes so far described, including hMSH2, hMLH1, hPMS1 and hPMS2 (Liu et al., Nature Genetics 9, 48, 1995). To ascertain whether 543X had a GTBP mutation, the RNA of this cell line was reverse transcribed with random hexamers and reverse transcriptase and the cDNA was then amplified with PCR using primers specific for the GTBP-coding sequence as reported above. In vitro transcription and translation of GTBP-coding sequence from 543X revealed truncated GTBP polypeptide after electrophoretic analysis. The sequence of the DNA region encoding GTBP was found to contain a 1 bp insertion (a T) at nucleotide 1876 of SEQ ID NO: 12, resulting in a frameshift and a downstream termination codon. The same mutation was identified also in the tumor tissue from this patient, but not in normal colon tissue. This proves that the mutation was somatic in nature and that it did not occur after the establishment of the cell culture line.

SEQUENCE LISTING

			SEQUENCE LISTING
			GENERAL INFORMATION
		(i)	APPLICANT: ISTITUTO DI RICERCHE DI BIOLOGIA
			MOLECOLARE P. ANGELETTI S.p.A.
5	1	(ii)	TITLE OF INVENTION: POLYPEPTIDE FOR
			REPAIRING GENETIC INFORMATION, NUCLEOTIDIC
			SEQUENCE WHICH CODES FOR IT AND PROCESS
			FOR THE PREPARATION THEREOF
	•	(iii)	NUMBER OF SEQUENCES: 16
10		(iv)	CORRESPONDENCE ADDRESS:
			(A) ADDRESSEE: Societa Italiana Brevetti
			(B)STREET: Piazza di Pietra, 39
			(C) CITY: Rome
			(D) COUNTRY: Italy
15			(E) POSTAL CODE: 1-00186
		(v)	COMPUTER READABLE FORM:
			(A) MEDIUM TYPE: Floppy disk 3.5" 1.44
			MBYTES
			(B) COMPUTER: IBM PC compatible
20			(C)OPERATING SYSTEM: PC-DOS/MS-DOS Rev.5.0
			(D)SOFTWARE: Microsoft Word 6.0
		(viii)	ATTORNEY INFORMATION
			(A) NAME: DI CERBO, Mario (Dr.)
			(C) REFERENCE: RM/X88551/PC-DC
25		(ix)	TELECOMMUNICATION INFORMATION
			(A) TELEPHONE: 06/6785941
			(B) TELEFAX: 06/6794692
			(C) TELEX: 612287 ROPAT
30	(1)	INFORM	ATION FOR SEQ ID NO: 1:
		(i)	SEQUENCE CHARACTERISTICS
			(A) LENGTH: 1292 amino acids
			(B) TYPE: amino acid
			(C)STRANDEDNESS: single
35			(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(iii) HYPOTHETICAL: No

			(iv)	ANT	ISE	NSE :	:	N)							
			(vi)	ORI	GIN	AL S	SOUR	.CE:								
					(A)	ORG	ANIS	: MS	Hot	mo s	sapi	ens					
			(vi	i)	IMM	EDI	ATE	SOU	RCE	: cI	ONA	clo	ne p	CIT	E2b	-C1	
5			(ix))	FEA	TURI	Ξ: S	EQ	ID 1	: 07	1 s	hows	s th	ne 1	292	amin	.0
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					enc	odeo	d by	r cl	one	Cl	(se	e SI	EQ I	D N	0: 3	12).	The
					sev	en d	olig	ope	ptic	les	whi	ch v	vere	id	ent:	ified	
					upo	n pı	rote	oly	tic	cle	ava	ge d	of G	TBP	(se	ee SE	Q
10					ID 1	NO:	2 t	.0 8) ar	e u	nde:	rlir	ned.	The	e fi	irst	
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					the	FLY	?5 c	DNA	is	Asn	at	pos	siti	on :	116.	•	
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					(C)	IDE	TIF	'ICA	TIOI	1 ME	THO	D: H	Expe	rim	enta	ally	
15			(xi))	SEQ	UENC	CE D	ESC	RIPT	CION	: S	EQ I	ID N	10:	1:		
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	1				5					10					15		
	Pro	Thr	Ser	Cys	Asp	Phe	Ser	Pro	Gly	Asp	Leu	Val	Trp	Ala	Lys	Met	
				20					25					30			
20	Glu	Gly		Pro	Trp	Trp	Pro	Cys	Leu	Val	Tyr	Asn	His	Pro	Phe	Asp	
			35					40					45				
	GIA		Phe.	Ile	Arg	Glu		Gly	Lys	Ser	Val		Val	His	Val	Gln	
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25	65	РЛЕ	ASD	ASD	Ser		Thr	Arg	GIY	Trp		ser	Lys	Arg	Leu		
23		Dwo	TT-s read	The se	Gly	70	T + + -	C 0 30	T	a 3	75	a 1	T	a 1	0 1	80	
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	Phe	Tvr	Ser	Δla	Lys	Pro	Glu	Tle	T.en		Δla	Mot	Gl n	7) ***	95	Acn	
		-7-		100	2,0		014		105	9	AIG	Mec	GIII	110	ATO	rap	
30	Glu	Ala	Leu		Lys	Asp	Lvs	Ile		Ara	Len	Glu	T.e.1		Va 1	Cre	
			115					120	-4-	3			125	•••	****	Cyb	
	Asp	Glu		Ser	Glu	Pro	Glu		Glu	Glu	Glu	Met		Val	Glv	Thr	
	-	130					135					140		-	2		
	Thr	Tyr	Val	Thr	Asp	Lys	Ser	Glu	Glu	Asp	Asn	Glu	Ile	Glu	Ser	Glu	
35	145	•			_	150				=	155					160	
	Glu	Glu	Val	Gln	Pro	Lys	Thr	Gln	Gly	Ser	Arg	Arg	Ser	Ser	Arg		
					165				_	170	_	•			175		

-45-

	Ile	Lys	Lys	Arg	Arg	Val	Ile	Ser	Asp	Ser	Glu	Ser	Asp	Ile	Gly	Gly
				180					185					190		
	Ser	Asp	Val	Glu	Phe	Lys	Pro	Asp	Thr	Lys	Glu	Glu	Gly	Ser	Ser	Asp
			195					200					205			
5	Glu	Ile	Ser	Ser	Gly	Val	Gly	Asp	Ser	Glu	Ser	Glu	Gly	Leu	Asn	Ser
		210					215					220				
	Pro	Val	Lys	Val	Ala	Arg	Lys	Arg	Lys	Arg	Met	Val	Thr	Gly	Asn	Gly
	225					230					235					240
	Ser	Leu	Lys	Arg	Lys	Ser	Ser	Arg	Lys	Glu	Thr	Pro	Ser	Ala	Thr	Lys
10					245					250					255	
	Gln	Ala	Thr	Ser	Ile	Ser	Ser	Glu	Thr	Lys	Asn	Thr	Leu	Arg	Ala	Phe
				260					265					270		
	Ser	Ala	Pro	Gln	Asn	Ser	Glu	Ser	Gln	Ala	His	Val	Ser	Gly	Gly	Gly
			275					280					285			
15	Asp	Asp	Ser	Ser	Arg	Pro	Thr	Val	Trp	Tyr	His	Glu	Thr	Leu	Glu	Trp
		290					295					300				
	Leu	Lys	Glu	Glu	Lys	Arg	Arg	Asp	Glu	His	Arg	Arg	Arg	Pro	Asp	His
	305					310					315					320
	Pro	Asp	Phe	Asp	Ala	Ser	Thr	Leu	Tyr	Val	Pro	Glu	Asp	Phe	Leu	Asn
20					325					330					335	
	Ser	Cys	Thr	Pro	Gly	Met	Arg	Lys	Trp	Trp	Gln	Ile	Lys	Ser	Gln	Asn
				340					345					350		
	Phe	Asp	Leu	Val	Ile	Cys	Tyr	Lys	Val	Gly	Lys	Phe	Tyr	Glu	Leu	Tyr
			355					360					365			
25	His	Met	Asp	Ala	Leu	Ile	Gly	Val	Ser	Glu	Leu	Gly	Leu	Val	Phe	Met
		370					375					380				
	Lys	Gly	Asn	Trp	Ala	His	Ser	Gly	Phe	Pro		Ile	Ala	Phe	Gly	Arg
	385					390					395					400
	Tyr	Ser	Asp	Ser		Val	Gln	Lys	Gly	Tyr	Lys	Val	Ala	Arg		Glu
30					405					410					415	
	Gln	Thr	Glu		Pro	Glu	Met	Met	Glu	Ala	Arg	Cys	Arg	Lys	Met	Ala
				420					425					430		
	His	Ile	Ser	Lys	Tyr	Asp	Arg	Val	Val	Arg	Arg	Glu	Ile	Cys	Arg	Ile
			435					440					445			
35	Ile	Thr	Lys	Gly	Thr	Gln		Tyr	Ser	Val	Leu		Gly	Asp	Pro	Ser
		450					455					460				
	Glu	Asn	Tyr	Ser	Lys	Tyr	Leu	Leu	Ser	Leu	Lys	Glu	Lys	Glu	Glu	Asp

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	465					470					475					480
	Ser	Ser	Gly	His	Thr	Arg	Ala	Tyr	Gly	Val	Cys	Phe	Val	Asp	Thr	Ser
					485					490					495	
	Leu	Gly	Lys	Phe	Phe	Ile	Gly	Gln	Phe	Ser	Asp	Asp	Arg	His	Cys	Ser
5				500					505					510		
	Arg	Phe	Arg	Thr	Leu	Val	Ala	His	Tyr	Pro	Pro	Val	Gln	Val	Leu	Phe
			515					520					525			
	Glu	Lys	Gly	Asn	Leu	Ser	Lys	Glu	Thr	Lys	Thr	Ile	Leu	Lys	Ser	Ser
		530					535					540				
10	Leu	Ser	Cys	Ser	Leu	Gln	Glu	Gly	Leu	Ile	Pro	Gly	Ser	Gln	Phe	Trp
	545					550					555					560
	Asp	Ala	Ser	Lys	Thr	Leu	Arg	Thr	Leu	Leu	Glu	Glu	Glu	Tyr	Phe	Arc
					565					570					575	
	Glu	Lys	Leu	Ser	Asp	Gly	Ile	Gly	Val	Met	Leu	Pro	Gln	Val	Leu	Lys
15				580					585					590		
	Gly	Met	Thr	Ser	Glu	Ser	Asp	Ser	Ile	Gly	Leu	Thr	Pro	Gly	Glu	Lys
			595					600					605			
	Ser	Glu	Leu	Ala	Leu	Ser	Ala	Leu	Gly	Gly	Cys	Val	Phe	Tyr	Leu	Lys
		610					615					620				
20	Lys	Cys	Leu	Ile	Asp		Glu	Leu	Leu	Ser	Met	Ala	Asn	Phe	Glu	Glu
	625		٠			630					635					640
	Tyr	Ile	Pro	Leu	Asp	Ser	Asp	Thr	Val	Ser	Thr	Thr	Arg	Ser	Gly	Ala
					645		_			650					655	
	Ile	Phe	Thr		Ala	Tyr	Gln	Arg		Val	Leu	Asp	Ala		Thr	Leu
25				660			_	_	665					670	_	_
	Asn	Asn	Leu	Glu	Ile	Pne	Leu		GIÀ	Thr	Asn	Gly		Thr	Glu	Gly
		_	675 -		_		_	680	_		_,	_	685		_	
	Thr		Leu	Glu	Arg	Val		Thr	Cys	His	Thr		Phe	GIA	Lys	Arg
	_	690	_		_	_	695		_	_	_	700				
30		Leu	Lys	Gin	Trp		Cys	Ala	Pro	Leu		Asn	His	Tyr	Ala	
	705	•	•			710	T 1 -	G3	3	•	715	**- 1		_	_	720
	Asn	Asp	Arg	Leu		Ата	TTE	GIU	Asp		Met	vaı	Val	Pro		Lys
	7 1.	a	~ 3	**- 1	725	61	T	7	T	730	•	~	_	_	735	_
35	тте	ser	Glu		vai	GIU	ьeu	ьеп		LYS	ьeu	Pro	ASD		GIU	Aro
, ,	T	T	C	740	- 7 -	TT	7 ~~	17-1	745	C	D	T	7	750	~ 12	3
	ren	Ten	Ser	шys	тте	UTS	ASII	760	GTÅ	ser	Pro	ren	Lys	ser	GIN	AST
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	HIS	PIO	Asp	Ser	Arg	Ата	TTE	Mec	Tyr	GIU	GIU	THE	THE	TYE	ser	гÀг
		770					775					780				
	Lys	Lys	Ile	Ile	Asp	Phe	Leu	Ser	Ala	Leu	Glu	Gly	Phe	Lys	Val	Met
	785					790		•			795					800
5	Cys	Lys	Ile	Ile	Gly	Ile	Met	Glu	Glu	Val	Ala	Asp	Gly	Phe	Lys	Ser
					805					810					815	
	Lys	Ile	Leu	Lys	Gln	Val	Ile	Ser	Leu	Gln	Thr	Lys	Asn	Pro	Glu	Gly
				820					825					830		
	Arg	Phe	Pro	Asp	Leu	Thr	Val	Glu	Leu	Asn	Arg	Trp	Asp	Thr	Ala	Phe
10			835					840					845			
	Asp	His	Glu	Lys	Ala	Arg	Lys	Thr	Gly	Leu	Ile	Thr	Pro	Lys	Ala	Gly
		850					855					860				
	Phe	Asp	Ser	Asp	Tyr	Asp	Gln	Ala	Leu	Ala	Asp	Ile	Arg	Glu	Asn	Glu
	865					870					875		Glu	Asn		
15					1045	5				1050)				1055	5
	Gly	Lys	Ala	Tyr	Cys	Val	Leu	Val	Thr	Gly	Pro	Asn	Met	Gly	Gly	Lys
				1060)				1065	5				1070)	
	Ser	Thr	Leu	Met	Arg	Gln	Ala	Gly	Leu	Leu	Ala	Val	Met	Ala	Gln	Met
			1075	5				1080)				1085	5		
20	Gly	Cys	Tyr	Val	Pro	Ala	Glu	Val	Cys	Arg	Leu	Thr	Pro	Ile	Asp	Arg
		1090)				1095	5				1100)			
	Val	Phe	Thr	Arg	Leu	Gly	Ala	Ser	Asp	Arg	Ile	Met	Ser	Gly	Glu	Ser
	1109	5				1110)				1115	5				1120
	Thr	Phe	Phe	Val	Glu	Leu	Ser	Glu	Thr	Ala	Ser	Ile	Leu	Met	His	Ala
25					1125	5				1130)				1135	5
	Thr	Ala	His	Ser	Leu	Val	Leu	Val	Asp	Glu	Leu	Gly	Arg	Gly	Thr	Ala
				1140	D				1145	5				115	כ	
	Thr	Phe	Asp	Gly	Thr	Ala	Ile	Ala	Asn	Ala	Val	Val	Lys	Glu	Leu	Ala
			1155	5				1160)				116	5		
30	Glu	Thr	Ile	Lys	Cys	Arg	Thr	Leu	Phe	Ser	Thr	His	Tyr	His	Ser	Leu
		117	0				117	5				118	0			
	Val	Glu	Asp	Tyr	Ser	Gln	Asn	Val	Ala	Val	Arg	Leu	Gly	His	Met	Ala
	118	5				119	0				119	5				1200
	Cys	Met	Val	Glu	Asn	Glu	Cys	Glu	Asp	Pro	Ser	Gln	Glu	Thr	Ile	Thr
35					120	5				121	0				121	5
	Phe	Leu	Tyr	Lys	Phe	Ile	Lys	Gly	Ala	Cys	Pro	Lys	Ser	Tyr	Gly	Phe
				122	0				122	5				123	0,	

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Asn Ala Ala Arg Leu Ala Asn Leu Pro Glu Glu Val Ile Gln Lys Gly 1235 1240 1245 His Arg Lys Ala Arg Glu Phe Glu Lys Met Asn Gln Ser Leu Arg Leu 1255 5 Phe Arg Glu Val Cys Leu Ala Ser Glu Arg Ser Thr Val Asp Ala Glu 1270 1275 1280 Ala Val His Lys Leu Leu Thr Leu Ile Lys Glu Leu 1285 1290 (2) INFORMATION FOR SEQ ID NO: 2: 10 (i) SEQUENCE CHARACTERISTICS (A) LENGTH: 10 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: protein (iii) HYPOTHETICAL: No (iv) ANTISENSE: No (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo sapiens 20 (ix) FEATURE: SEQ ID NO: 2 to 8 show seven oligopeptides derived from proteolytic cleavage of GTBP extracted from HeLa cells and purified as described in ref. 16 . The peptide corresponding to SEQ ID NO: 6 (18 amino acids) was selected to design two 25 degenerate primers corresponding to the Nand C-terminal sequences of the peptide, as given in detail in SEQ ID NO: 9 and 10. (A) NAME: FR44 30 (C) IDENTIFICATION METHOD: Experimentally SEQUENCE DESCRIPTION: SEQ ID NO: 2: Val Arg Val His Val Gln Phe Phe Asp Asp 10 (3) INFORMATION FOR SEQ ID NO: 3: 35 (i) SEQUENCE CHARACTERISTICS (A) LENGTH: 18 amino acids (B) TYPE: amino acid

		(C)STRANDEDNESS: single	
		(D) TOPOLOGY: linear	
	(ii	i) MOLECULE TYPE: protein	
	(ii	ii) HYPOTHETICAL: No	
5	(iv	v) ANTISENSE: No	
	(vi	i) ORIGINAL SOURCE:	
		(A) ORGANISM: Homo sapiens	
	(ix	x) FEATURE: see SEQ ID NO: 2	
		(A) NAME: FR48	
10		(C) IDENTIFICATION METHOD: Experimentally	
	(xi	i) SEQUENCE DESCRIPTION: SEQ ID NO: 3:	
	Lys Leu Pro	o Asp Leu Glu Arg Leu Leu Ser Lys Ile His Asn Val XXX	
	1	5 10	15
	Ser Lys		
15	(4) INF	FORMATION FOR SEQ ID NO: 4:	
	(i)	SEQUENCE CHARACTERISTICS	
		(A) LENGTH: 13 amino acids	
		(B) TYPE: amino acid	
		(C)STRANDEDNESS: single	
20		(D) TOPOLOGY: linear	
	(ii	i) MOLECULE TYPE: protein	
	(ii	ii) HYPOTHETICAL: No	
	(iv	v) ANTISENSE: No	
	(vi	i) ORIGINAL SOURCE:	
25		(A)ORGANISM: Homo sapiens	
	(ix	x) FEATURE: see SEQ ID NO: 2	
		(A) NAME: FR49b	
		(C) IDENTIFICATION METHOD: Experimentally	
	(xi	i) SEQUENCE DESCRIPTION: SEQ ID NO: 4:	
30	Leu Ser Arg	g Gly Iso Gly Val Met Leu Pro Gln Val Leu	
	1	5 10	
	(5) INF	FORMATION FOR SEQ ID NO: 5:	
	(i)) SEQUENCE CHARACTERISTICS	
		(A) LENGTH: 14 amino acids	
35		(B) TYPE: amino acid	
		(C)STRANDEDNESS: single	
		(D) TOPOLOGY: linear	

	(ii)	MOLECULE TYPE: protein
	(iii)	HYPOTHETICAL: No
	(iv)	ANTISENSE: No
	(vi)	ORIGINAL SOURCE:
5		(A)ORGANISM: Homo sapiens
	(ix)	FEATURE: see SEQ ID NO: 2
		(A) NAME: FR49c
		(C) IDENTIFICATION METHOD: Experimentally
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 5:
10	Thr Leu Arg Th	ur Leu Leu Glu Glu Tyr Phe Arg Glu Lys
	1	5 10
	(6) INFORM	MATION FOR SEQ ID NO: 6:
	(i)	SEQUENCE CHARACTERISTICS
		(A) LENGTH: 18 amino acids
15		(B) TYPE: amino acid
		(C)STRANDEDNESS: single
	•	(D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: protein
	(iii)	HYPOTHETICAL: No
20	(iv)	ANTISENSE: No
	(vi)	ORIGINAL SOURCE:
		(A)ORGANISM: HeLa cell extract
	(ix)	FEATURE: see SEQ ID NO: 2
		(A) NAME: FR52
25		(C) IDENTIFICATION METHOD: Experimentally
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 6:
	Ser Tyr Gly Ph	e Asn Ala Ala Arg Leu Ala Asn Leu Pro Glu Glu Val
	1	5 10 15
	Ile Gln	
30		MATION FOR SEQ ID NO: 7:
	(i)	SEQUENCE CHARACTERISTICS
		(A) LENGTH: 13 amino acids
		(B) TYPE: amino acid
		(C)STRANDEDNESS: single
35		(D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: protein

		(iii)	HYPOTHETICAL: No
		(iv)	ANTISENSE: No
		(vi)	ORIGINAL SOURCE:
			(A)ORGANISM: Homo sapiens
5		(ix)	FEATURE: see SEQ ID NO: 2
			(A) NAME: FR59
			(C) IDENTIFICATION METHOD: Experimentally
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 7:
	Asn Pr	o Glu Gly	y Arg Phe Pro Asp Leu Thr Val Glu Leu
10	1		5 10
	(8)	INFORM	ATION FOR SEQ ID NO: 8:
		(i)	SEQUENCE CHARACTERISTICS
			(A) LENGTH: 11 amino acids
			(B) TYPE: amino acid
15			(C)STRANDEDNESS: single
			(D) TOPOLOGY: linear
		(ii)	MOLECULE TYPE: protein
		(iii)	HYPOTHETICAL: No
		(iv)	ANTISENSE: No
20		(vi)	ORIGINAL SOURCE:
			(A)ORGANISM: Homo sapiens
		(ix)	FEATURE: see SEQ ID NO: 2
			(A) NAME: FR69
			(C) IDENTIFICATION METHOD: Experimentally
25		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 8:
	Ile I	le Asp	Phe Leu Ser Ala Leu Glu Gly Phe
	1		5 10
	(9)	INFORM	ATION FOR SEQ ID NO: 9
		(i)	SEQUENCE CHARACTERISTICS
30			(A) LENGTH: 22 base pairs
			(B) TYPE: nucleic acid
			(C)STRANDEDNESS: single
			(D) TOPOLOGY: linear
		(ii)	MOLECULE TYPE: synthetic DNA
35			HYPOTHETICAL: No
			ANTISENSE: No
		(vii)	IMMEDIATE SOURCE: oligonucleotide synthesizer

	(ix)	FEATURE: SEQ ID NO:9 shows the sequence of
		the degenerate single-stranded DNA primer
		deduced from the N-terminal of oligopeptide
		shown in SEQ ID NO: 6. Together with SEQ ID
5		NO: 10, the two primers were used to amplify
		poly-A ⁺ RNA extracted from HeLa cells. The
		expected 67 base pairs (bp) fragment was
		cloned in pBluescript SK- (Stratagene) and
		sequenced with a commercial T7-polymerase
10		based kit (Pharmacia). The 54 bp sequence of
		the resulting fragment, obtained after
		subtraction of the engineered cloning sites,
		is shown as SEQ ID NO: 11.
		(A) NAME: oligo 5' sense
15		(C) IDENTIFICATION METHOD: Polyacrylamide gel
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 9
	GCGAATTCTA YGG	NTTYAAY GC 22
	(10) INFORM	ATION FOR SEQ ID NO: 10
	(i)	SEQUENCE CHARACTERISTICS
20		(A)LENGTH: 22 base pairs
		(B) TYPE: nucleic acid
		(C)STRANDEDNESS: single
		(D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: synthetic DNA
25	(iii)	HYPOTHETICAL: No
	(iv)	ANTISENSE: Yes
	(vii)	IMMEDIATE SOURCE: oligonucleotide synthesizer
	(ix)	FEATURE: SEQ ID NO:10 shows the sequence of the
		degenerate single-stranded DNA primer deduced
30		from the C-terminal of oligopeptide shown in SEQ
		ID NO: 6. Together with SEQ ID NO: 9, the two
		primers were used to amplify poly-A ⁺ RNA
		extracted from HeLa cells. The expected 67 base
		pairs (bp) fragment was cloned in pBluescript
35		SK- (Stratagene) and sequenced with a commercial
		T7-polymerase based kit (Pharmacia). The 54 bp

sequence of the resulting fragment, obtained

10

15

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		after subtraction of the engineered cloning
		sites, is shown as SEQ ID NO: 11.
		(A)NAME: oligo 3' antisense
		(C) IDENTIFICATION METHOD: Polyacrylamide gel
	(xi)	_
GCGGAT	CCTC YTG	DATNACY TC 22
(11)	INFORM	MATION FOR SEQ ID NO: 11
	(i)	SEQUENCE CHARACTERISTICS
		(A) LENGTH: 54 base pairs
		(B) TYPE: nucleic acid
		(C)STRANDEDNESS: double
		(D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: synthetic DNA
	(iii)	HYPOTHETICAL: No
	(iv)	ANTISENSE: Yes
	(vii)	IMMEDIATE SOURCE: PCR product
	(ix)	FEATURE: SEQ ID NO: 11 shows the double-stranded
		DNA sequence encoding the oligopeptide reported
		in SEQ ID NO: 6, as deduced by sequencing of
		cloned amplification product. This fragment was
		derived from PCR amplification of HeLa cDNA,
		using the degenerate primers described in SEQ ID
		NO: 9 and 10. The DNA sequence was end-labelled
		with ³² P by a standard kinase reaction (with T4
		polynucleotide kinase PNK and [g-32P]ATP as
		described by Maniatis et al., Molecular cloning:
		a laboratory manual, Cold Spring Harbor
		Laboratory, Cold Spring Harbor, N.Y., 1982) in
		order to generate a double-stranded DNA probe.
		The labelled probe was used in the screening of
		a commercial oligo dT-primed cDNA library in
		phage lambda (HeLa S3 UNI-ZAP XR, Stratagene).
		Screening of the HeLa S3 UNI-ZAP XR library in
		phage lambda made it possible the identification
		of two clones hybridizing with the DNA probe.
		or one crosses injurically with the DNA probe.

These clones were designated C1 and FLY5.

(A) NAME:

			(C) ID	ENTIFICAT	TON METHO	D: Polyac	rylamide -	gel
	(xi)	SEQUE	NCE DESCR	RIPTION: S	EQ ID NO:	11	
	AGCTATGG	CT TTAA	TGCAGC	AAGGCTTGCT	· AATCTCCCAG	AGGAAGTTAT	TCAA	
	54							
5	(12) I	NFORM	NOIT	FOR SEQ	D NO: 12			
	(i)	SEQUE	NCE CHARA	CTERISTIC	S		
			(A) LE	NGTH: 398	0 base pa	irs		
			(B)TY	PE: nucle	ic acid			
			(C)ST	RANDEDNES	S: double			
10			(D) TO	POLOGY: 1	inear			
	(ii)	MOLEC	ULE TYPE:	syntheti	c DNA		
	(iii)	HYPOT	HETICAL:	No			
	(iv)	ANTIS	ENSE: No				
	(vii)	IMMED	IATE SOUR	CE: cDNA	clone C1		
15	(ix)	FEATU.	RE: SEQ I	D NO: 12	shows the	3980 bp (CDNA
			seque	nce of cl	one C1. T	he cDNA i	nsert of o	clone
			FLY5	spanned f	rom nucle	otide 346	to 3980 d	of the
			C1 se	quence as	reported	in SEQ I	D NO: 12.	
			(A) NA	ME: C1				
20			(C) ID	ENTIFICAT	ION METHO	D: Polyac:	rylamide g	gel
	(:	xi)	SEQUE	NCE DESCR	IPTION: S	EQ ID NO:	12	
	GCGAAGAA	CC TCAA	CGGAGG	GCTGCGGAGA	TCGGTAGCGC	CTGCTGCCCC	CACCAGTTGT	60
	GACTTCTC	AC CAGG	AGATTT	GGTTTGGGCC	AAGATGGAGG	GTTACCCCTG	GTGGCCTTGT	120
	CTGGTTTA	CA ACCA	CCCCTT	TGATGGAACA	TTCATCCGCG	AGAAAGGGAA	ATCAGTCCGT	180
25	GTTCATGT	AC AGTT	TTTTGA	TGACAGCCCA	ACAAGGGGCT	GGGTTAGCAA	AAGGCTTTTA	240
	AAGCCATAT	ra cagg:	TTCAAA	ATCAAAGGAA	GCCCAGAAGG	GAGGTCATTT	TTACAGTGCA	300
	AAGCCTGAA	AA TACTO	GAGAGC	AATGCAACGT	GCAGATGAAG	CCTTAAATAA	AGACAAGATT	360
	AAGAGGCTT	G AATT	GCAGT	TTGTGATGAG	CCCTCAGAGC	CAGAAGAGGA	AGAAGAGATG	420
	GAGGTAGG	CA CAAC	TTACGT	AACAGATAAG	AGTGAAGAAG	ATAATGAAAT	TGAGAGTGAA	480
30	GAGGAAGTA	AC AGCC	TAAGAC	ACAAGGATCT	AGGCGAAGTA	GCCGCCAAAT	AAAAAAACGA	540
	AGGGTCATA	T CAGA	TTCTGA	GAGTGACATT	GGTGGCTCTG	ATGTGGAATT	TAAGCCAGAC	600
	ACTAAGGAG	G AAGG	AAGCAG	TGATGAAATA	AGCAGTGGAG	TGGGGGATAG	TGAGAGTGAA	660
	GGCCTGAAC	CA GCCC	rgtcaa	AGTTGCTCGA	AAGCGGAAGA	GAATGGTGAC	TGGAAATGGC	720
	TCTCTTAAA	AA GGAAA	AAGCTC	TAGGAAGGAA	ACGCCCTCAG	CCACCAAACA	AGCAACTAGC	780
35	ATTTCATCA	G AAAC	CAAGAA	TACTTTGAGA	GCTTTCTCTG	CCCCTCAAAA	TTCTGAATCC	840
	CAAGCCCAC	G TTAG	rggagg	TGGTGATGAC	AGTAGTCGCC	CTACTGTTTG	GTATCATGAA	900
	א כייייייא כיא א	יד מממייי	רא א כיכיא	CCDDDDCDCD	A CA CA TCA CC	20202022	GGGMG3 = G3 G	0.00

	CCCGATTTTG	ATGCATCTAC	ACTCTATGTG	CCTGAGGATT	TCCTCAATTC	TTGTACTCCT	1020
	GGGATGAGGA	AGTGGTGGCA	GATTAAGTCT	CAGAACTTTG	ATCTTGTCAT	CTGTTACAAG	1080
	GTGGGGAAAT	TTTATGAGCT	GTACCACATG	GATGCTCTTA	TTGGAGTCAG	TGAACTGGGG	1140
	CTGGTATTCA	TGAAAGGCAA	CTGGGCCCAT	TCTGGCTTTC	CTGAAATTGC	ATTTGGCCGT	1200
5	TATTCAGATT	CCCTGGTGCA	GAAGGGCTAT	AAAGTAGCAC	GAGTGGAACA	GACTGAGACT	1260
	CCAGAAATGA	TGGAGGCACG	ATGTAGAAAG	ATGGCACATA	TATCCAAGTA	TGATAGAGTG	1320
	GTGAGGAGGG	AGATCTGTAG	GATCATTACC	AAGGGTACAC	AGACTTACAG	TGTGCTGGAA	1380
	GGTGATCCCT	CTGAGAACTA	CAGTAAGTAT	CTTCTTAGCC	TCAAAGAAAA	AGAGGAAGAT	1440
	TCTTCTGGCC	ATACTCGTGC	ATATGGTGTG	TGCTTTGTTG	ATACTTCACT	GGGAAAGTTT	1500
10	TTCATAGGTC	AGTTTTCAGA	TGATCGCCAT	TGTTCGAGAT	TTAGGACTCT	AGTGGCACAC	1560
	TATCCCCCAG	TACAAGTTTT	ATTTGAAAAA	GGAAATCTCT	CAAAGGAAAC	TAAAACAATT	1620
	CTAAAGAGTT	CATTGTCCTG	TTCTCTTCAG	GAAGGTCTGA	TACCCGGCTC	CCAGTTTTGG	1680
	GATGCATCCA	AAACTTTGAG	AACTCTCCTT	GAGGAAGAAT	ATTTTAGGGA	AAAGCTAAGT	1740
	GATGGCATTG	GGGTGATGTT	ACCCCAGGTG	CTTAAAGGTA	TGACTTCAGA	GTCTGATTCC	1800
15	ATTGGGTTGA	CACCAGGAGA	GAAAAGTGAA	TTGGCCCTCT	CTGCTCTAGG	TGGTTGTGTC	1860
	TTCTACCTCA	AAAAATGCCT	TATTGATCAG	GAGCTTTTAT	CAATGGCTAA	TTTTGAAGAA	1920
	TATATTCCCT	TGGATTCTGA	CACAGTCAGC	ACTACAAGAT	CTGGTGCTAT	CTTCACCAAA	1980
	GCCTATCAAC	GAATGGTGCT	AGATGCAGTG	ACATTAAACA	ACTTGGAGAT	TTTTCTGAAT	2040
	GGAACAAATG	GTTCTACTGA	AGGAACCCTA	CTAGAGAGGG	TTGATACTTG	CCATACTCCT	2100
20	TTTGGTAAGC	GGCTCCTAAA	GCAATGGCTT	TGTGCCCCAC	TCTGTAACCA	TTATGCTATT	2160
	AATGATCGTC	TAGATGCCAT	AGAAGACCTC	ATGGTTGTGC	CTGACAAAAT	CTCCGAAGTT	2220
	GTAGAGCTTC	TAAAGAAGCT	TCCAGATCTT	GAGAGGCTAC	TCAGTAAAAT	TCATAATGTT	2280
	GGGTCTCCCC	TGAAGAGTCA	GAACCACCCA	GACAGCAGGG	CTATAATGTA	TGAAGAAACT	2340
	ACATACAGCA	AGAAGAAGAT	TATTGATTTT	CTTTCTGCTC	TGGAAGGATT	CAAAGTAATG	2400
25	TGTAAAATTA	TAGGGATCAT	GGAAGAAGTT	GCTGATGGTT	TTAAGTCTAA	AATCCTTAAG	2460
	CAGGTCATCT	CTCTGCAGAC	AAAAAATCCT	GAAGGTCGTT	TTCCTGATTT	GACTGTAGAA	2520
	TTGAACCGAT	GGGATACAGC	CTTTGACCAT	GAAAAGGCTC	GAAAGACTGG	ACTTATTACT	2580
	CCCAAAGCAG	GCTTTGACTC	TGATTATGAC	CAAGCTCTTG	CTGACATAAG	AGAAAATGAA	2640
	CAGAGCCTCC	TGGAATACCT	AGAGAAACAG	CGCAACAGAA	TTGGCTGTAG	GACCATAGTC	2700
30	TATTGGGGGA	TTGGTAGGAA	CCGTTACCAG	CTGGAAATTC	CTGAGAATTT	CACCACTCGC	2760
	AATTTGCCAG	AAGAATACGA	GTTGAAATCT	ACCAAGAAGG	GCTGTAAACG	ATACTGGACC	2820
	AAAACTATTG	AAAAGAAGTT	GGCTAATCTC	ATAAATGCTG	AAGAACGGAG	GGATGTATCA	2880
	TTGAAGGACT	GCATGCGGCG	ACTGTTCTAT	AACTTTGATA	AAAATTACAA	GGACTGGCAG	2940
	TCTGCTGTAG	AGTGTATCGC	AGTGTTGGAT	GTTTTACTGT	GCCTGGCTAA	CTATAGTCGA	3000
35	GGGGGTGATG	GTCCTATGTG	TCGCCCAGTA	ATTCTGTTGC	CGGAAGATAC	CCCCCCTTC	3060
	TTAGAGCTTA	AAGGATCACG	CCATCCTTGC	ATTACGAAGA	CTTTTTTTGG	AGATGATTTT	3120
	ATTCCTAATG	ACATTCTAAT	AGGCTGTGAG	GAAGAGGAGC	AGGAAAATGG	CAAAGCCTAT	3180

	TGTGTGCTTG TTACTGGACC AAATATGGGG GGCAAGTCTA CGCTTATGAG ACAGGCTGGC	3240
	TTATTAGCTG TAATGGCCCA GATGGGTTGT TACGTCCCTG CTGAAGTGTG CAGGCTCACA	3300
	CCAATTGATA GAGTGTTTAC TAGACTTGGT GCCTCAGACA GAATAATGTC AGGTGAAAGT	3360
	ACATTTTTG TTGAATTAAG TGAAACTGCC AGCATACTCA TGCATGCAAC AGCACATTCT	3420
5	CTGGTGCTTG TGGATGAATT AGGAAGAGGT ACTGCAACAT TTGATGGGAC GGCAATAGCA	3480
	AATGCAGTTG TTAAAGAACT TGCTGAGACT ATAAAATGTC GTACATTATT TTCAACTCAC	3540
	TACCATTCAT TAGTAGAAGA TTATTCTCAA AATGTTGCTG TGCGCCTAGG ACATATGGCA	3600
	TGCATGGTAG AAAATGAATG TGAAGACCCC AGCCAGGAGA CTATTACGTT CCTCTATAAA	3660
	TTCATTAAGG GAGCTTGTCC TAAAAGCTAT GGCTTTAATG CAGCAAGGCT TGCTAATCTC	3720
10	CCAGAGGAAG TTATTCAAAA GGGACATAGA AAAGCAAGAG AATTTGAGAA GATGAATCAG	3780
	TCACTACGAT TATTTCGGGA AGTTTGCCTG GCTAGTGAAA GGTCAACTGT AGATGCTGAA	3840
	GCTGTCCATA AATTGCTGAC TTTGATTAAG GAATTATAGA CTGACTACAT TGGAAGCTTT	3900
	GAGTTGACTT CTGACCAAAG GTGGTAAATT CAGACAACAT TATGATCTAA TAAACTTTAT	3960
	TTTTTAAAA TGAAAAAAA	
15	3980	
	(13) INFORMATION FOR SEQ ID NO: 13	
	(i) SEQUENCE CHARACTERISTICS	
	(A) LENGTH: 394 base pairs	
	(B) TYPE: nucleic acid	
20	(C)STRANDEDNESS: double	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: synthetic DNA	
	(iii) HYPOTHETICAL: No	
	(iv) ANTISENSE: No	
25 .	(vii) IMMEDIATE SOURCE: Homo sapiens	
	(ix) FEATURE: SEQ ID NO: 13 shows the double-stra	nded
	DNA sequence used to express an internal dor	nain
	of hMSH2 (corresponding to amino acid residu	
	27 to 158) in the expression vector pGEX-3x	
30	also legend to Figure 2).	
	(A) NAME: GST/hMSH2	
	(C) IDENTIFICATION METHOD: Polyacrylamide gel	L
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13	
	GGAGAAGCCG ACCACCACAG TGCGCCTTTT CGACCGGGGC GACTTCTATA CGGCGCACGG	60
35	CGAGGACGCG CTGCTGGCCG CCCGGGAGGT GTTCAAGACC CAGGGGGTGA TCAAGTACAT	120
	GGGGCCGGCA GGAGCAAAGA ATCTGCAGAG TGTTGTGCTT AGTAAAATGA ATTTTGAATC	180
	TTTTGTAAAA GATCTTCTTC TGGTTCGTCA GTATAGAGTT GAAGTTTATA AGAATAGAGC	240

	TGGAAATAAG GCATCCAAGG AGAATGATTG GTATTTGGCA TATAAGGCTT CTCCTGGCAA	300
	TCTCTCTCAG TTTGAAGACA TTCTCTTTGG TAACAATGAT ATGTCAGCTT CCATTGGTGT	360
	TGTGGGTGTT AAAATGTCCG CAGTTGATGG CCAG	394
	(14) INFORMATION FOR SEQ ID NO: 14	
5	(i) SEQUENCE CHARACTERISTICS	
	(A) LENGTH: 534 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: double	
	(D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: synthetic DNA	
	(iii) HYPOTHETICAL: No	
	(iv) ANTISENSE: No	
	(vii) IMMEDIATE SOURCE:	
	(ix) FEATURE: SEQ ID NO: 14 shows the double-stra	anded
15	DNA sequence used to express an internal dom	nain
	of GTBP (corresponding to amino acid residue	es
	750 to 928) in the expression vector pGEX-33	c
	(see also legend to Figure 2).	
	(A) NAME: GST/GTBP	
20	(C) IDENTIFICATION METHOD: Polyacrylamide gel	L
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14	
	CTTGAGAGGC TACTCAGTAA AATTCATAAT GTTGGGTCTC CCCTGAAAGT CAGAACCACC	60
	CAGACAGCAG GGCTATAATG TATGAAGAAA CTACATACAG CAAGAAGAAG ATTATTGATT	120
	TTCTTTCTGC TCTGGAAGGA TTCAAAGTAA TGTGTAAAAT TATAGGGATC ATGGAAGAAG	180
25	TTGCTGATGG TTTTAAGTCT AAAATCCTTA AGCAGGTCAT CTCTCTGCAG ACAAAAAATC	240
	CTGAAGGTCG TTTTCCTGAT TTGACTGTAG AATTGAACCG ATGGGATACA GCCTTTGACC	300
	ATGAAAAGGC TCGAAAGACT GGACTTATTA CTCCCAAAGC AGGCTTTGAC TCTGATTATG	360
	ACCAAGCTCT TGCTGACATA AGAGAAAATG AACAGAGCCT CCTGGAATAC CTAGAGAAAC	420
	AGCGCAACAG AATTGGCTGT AGGACCATAG TCTATGGATT GGTAGGAACC GTTACGCAGC	480
30	TGGAAATTCC TGAGAATTTC ACCACTCGCA ATTTGCCAGA AGAATACGAG TTGA	534
	(15) INFORMATION FOR SEQ ID NO: 15	
	(i) SEQUENCE CHARACTERISTICS	
	(A) LENGTH: 68 amino acids	
	(B) TYPE: amino acid	
35	(C)STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: protein	

		(iii)	HYPOTHETICAL: No
		(iv)	ANTISENSE: No
		(vi)	ORIGINAL SOURCE:
			(A)ORGANISM: Homo sapiens
5		(vii)	IMMEDIATE SOURCE: cDNA of clone KMN
		(ix)	FEATURE: SEQ ID NO: 15 shows the amino-terminal
			sequence of 68 amino acids of GTBP encoded by
			the clone TASNR2A1 (see SEQ ID NO:16 for the
			corresponding nucleotide encoding sequence). The
10			amino acid sequence SEQ ID NO:15 (corresponding
			to residues 1-68) must be placed in front of the
			amino acid in position 1 of the sequence given
			in SEQ ID NO:1 (corresponding to 1292 residues)
			to obtain the complete GTBP sequence of 1360
15			amino acids.
			(A) NAME: KMN
			(C) IDENTIFICATION METHOD: experimental
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 15
	Met Se	r Arg Glr	ser Thr Leu Tyr Ser Phe Phe Pro Lys Ser Pro Ala
20	1		5 10 15
	Lys Se	r Asp Ala	Met Lys Ala Ser Ala Arg Ala Ser Arg Glu Gly Gly
		20	
	Arg Ala	a Ala Ala	Ala Pro Glu Ala Ser Pro Ser Pro Gly Gly Asp Ala
		35	40 45
25	Ala Ty	r Ser Glu	Ala Gly Pro Gly Pro Arg Pro Leu Ala Arg Ser Ala
	50		55 60
		o Pro Lys	
	65 (3.6	TMEODM	AUTON FOR GEO TO NO. 16
30	(16 (i)		ATION FOR SEQ ID NO: 16 CE CHARACTERISTICS
50	(_ /		
			(A)LENGTH: 204 base pairs (B)TYPE: nucleic acid
			(C) STRANDEDNESS: double
			(D) TOPOLOGY: linear
35		(ii)	MOLECULE TYPE: synthetic DNA
-			HYPOTHETICAL: No
			ANTISENSE: No

GCGCGATCCG CGTCACCGCC CAAG

	(vii)	IMMEDIATE SOURCE: cDNA of clone KMN
	(ix)	FEATURE: SEQ ID NO: 16 shows the double-stranded
		DNA sequence obtained using the RACE method
		(Rapid Anmplification cDNA Ends) used to
5		establish the 5'-terminal sequence of GTBP cDNA
		encoding the amino-terminal region of the
		protein GTBP as indicated in SEQ ID NO:15. The
		nucleotidic sequence SEQ ID NO:15 (corresponding
		to 204 residues) must be positioned in front of
10		the nucleotide in position 1 of the sequence
		given in SEQ ID NO:12 (corresponding to 3980
		residues) in order to obtain the complete GTBP-
		encoding sequence of 4080 nucleotides.
		(A) NAME: KMN
15		(C) IDENTIFICATION METHOD: Polyacrylamide gel
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 16
	ATGTCGCGAC AGAC	GCACCCT GTACAGCTTC TTCCCCAACT CTCCGGCGCT GAGTGATGCC 60
	AACAAGGCCT CGG	CCAGGGC CTCACGCGAA GGCGGCCGTG CCGCCGCTGC CCCCGAGGCC 120
	TCTCCTTCCC CAG	GCGGGAA TGCGGCCTGG AGCGAGGCTG GGCCTGGGCC CAGGCCCTTG 180

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CLAIMS

- 1. An isolated polypeptide, wherein said polypeptide comprises: (1) a first sequence corresponding to GTBP as set forth by combining the amino acid sequences set forth in SEQ ID NO: 15 and SEQ ID NO:1; a second sequence wherein said second sequence is a subsequence of said first sequences and is at least 4 amino acids; (3) a third sequence in which at least one amino acid is replaced by a different amino acID
- 10 2. The polypeptide of Claim 1 complexed to a second polypeptide.
 - 3. The polypeptide complex of Claim 2, wherein said second polypeptide is hMSH2.
 - 4. An isolated polypeptide according to claim 1, comprising the amino acid sequences from amino acid 1 to 68 of SEQ ID NO:15 and from amino acid 1 to 1292 of SEQ ID NO: 1, or in any case sequences within the combination of SEQ ID NO: 15 and SEQ ID NO:1, for example SEQ ID NO: 2 to SEQ ID NO:8).
- 5. An isolated DNA or RNA molecule, wherein said molecule comprises:
 - (1) a first sequence encoding GTBP as set forth by combining SEQ ID NO:16 and SEQ ID NO: 12;
 - (2) a second sequence, wherein said second sequence is a subsequence of said first sequence and is at least 10 nucleotides in length;
 - (3) a third sequence in which at least one nucleotide of said first or second sequence is replaced by a different nucleotide; or
- 30 (4) a fourth sequence complementary to any of said first second, or third sequences;

with the provisos that (1) if said molecule is an RNA molecule, U replaces T in said sequence of said molecule, (2) said third sequence is at least 95% identical to said first or second sequence, and (3) said second sequence is not present in hMSH2 cDNA.

6. The molecule of Claim 5, wherein said molecule comprises said first sequence.

- 7. The molecule of Claim 5, wherein said molecule comprises said second sequence.
- 5 8. The molecule of Claim 5, wherein said molecule comprises said third sequence.
 - 9. The molecule of Claim 5, wherein said molecule comprises a cDNA sequence.
 - 10. The molecule of Claim 5, wherein said molecule consists essentially of DNA encoding GTBP.

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- 11. The molecule of Claim 5, wherein the RNA or DNA encoding GTBP is naturally occuring.
- 12. An expression vector containing the molecule of Claim 5.
- 13. A cell transformed with the molecule of Claim 5.
 - 14. The cell of Claim 13, wherein said molecule is DNA and said DNA is arranged in operative association with an expression control sequence capable of directing replication and expression of said DNA.
 - 15. The cell according to Claim 13, wherein said cell is a eukaryotic or prokaryotic cell including animal, fungal or bacterial cell.
 - 16. A process for producing GTBP protein comprising culturing a cell of Claim 13 in a suitable culture medium and isolating said GTBP protein from said cell.
 - 17. A polypeptide made according to the process of Claim 16.
 - 18. A method for identifying agents which inhibit or enhance GTBP activity as detectable by in vitro multior dimeriation assays, DNA-binding assays and mismatch repair assays.
 - 19. A method of identifying GTBP-modulating agents, comprising:
- 35 (1) performing a heterodimerization that includes a GTBP polypeptide, hMSH2 and an agent, and (2)

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detecting whether the agent modulates heterodimerization.

- 20. The method of Claim 19, wherein the heteodimerization assay comprises an *in vitro* binding reaction.
- 21. A preparation of specific antibodies immunoreactive with GTBP and not substantially immunoreactive with other proteins unrelated to GTBP.
- 22. A method of purification of GTBP or GTBP-complexing molecules involving the use of specific antibodies of Claim 21.
- 23. A method of purification of GTBP or GTBP-complexing molecules based on specific interaction between GTBP and nucleic acid recognition sequences.
- 24. A method of detecting the presence of a genetic defect that has the potential of causing tumorigenesis in human, which comprises:

identifying a mutation of a *GTBP* gene of said human, wherein said mutation results in a *GTBP* gene sequence different from wild-type human GTBP-coding DNA sequence as set forth by combining SEQ ID NO:16 and SEQ ID NO: 12.

- 25. A method of detecting the presence of a genetic defect that causes cancer in a human, which comprises:
- identifying a mutation of a GTBP gene of said human, wherein said mutation provides a GTBP gene sequence different from human GTBP DNA sequence as set forth by combining SEQ ID NO:16 and SEQ ID NO: 12, that changes the sequence of a protein product of said GTBP gene, or that causes the GTBP product to be truncated or that results in said GTBP gene not being transcribed or translated.
- 26. A method of diagnosing or prognosing a neoplastic tissue of a human comprising:
- identifying the presence of a mutation of a GTBP gene or its expression product in said tissue of said human patient, wherein said mutation provides a GTBP

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gene sequence different from human *GTBP* DNA sequence as set forth by combining SEQ ID NO:12 and SEQ ID NO: 16, said alteration indicating neoplasia of the tissue.

- 27. The methods of Claims 24-26, wherein said mutations result in a change in the sequence of a protein product of said *GTBP* gene.
- 28. The methods of Claims 24-26, wherein said mutations result in said *GTBP* gene not being transcribed or translated.
- 29. The methods of Claims 24-26, wherein said mutations create stop codons in said GTBP gene.
 - 30. The methods of Claims 24-26, wherein said methods comprise Polymerase Chain Reaction (PCR) amplification of at least a segment of said *GTBP* gene.
 - 31. The methods of Claims 24-26, whereas said methods comprise identifying a change in a restriction site as a result of said mutation.
 - 32. The methods of Claim 24-26, wherein said methods comprise restriction fragment length polymorphism analysis, allele-specific oligonucleotide hybridization or nucleotide sequencing.
 - 33. The methods of Claims 24-26, wherein said methods classify said human as homozygous for said GTBP gene or for said mutated GTBP gene or heterozygous for said GTBP gene and said mutated GTBP gene.
 - 34. The methods of Claims 24-26 wherein the expression products are mRNA molecules.
 - 35. The methods of Claims 24-26 wherein the loss of wild-type GTBP coding sequence is detected by Nothern hybridization of mRNA molecules extracted from cells or tissues.
 - 36. The methods of Claims 24-26 wherein the loss of wild-type *GTBP* is detected by Southern hybridization of a GTBP DNA probe to genomic DNA of said human patient.
 - 37. The methods of Claims 24-26 wherein the loss of wild-type *GTBP* gene is detected by identifying a mismatch between nucleic acids including (1) mRNA molecules of

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said human patient and (2) a nucleic acid complementary to human wild-type GTBP coding sequence, when molecules 1 and 2 are hybridized with each other and form a duplex.

- 38. The methods of Claims 24-26 wherein the loss of wild-type gene is detected by gene cloning and sequencing of cloned DNA.
- 39. The methods of Claims 24-26 wherein the loss of wild-type *GTBP* gene is detected by screening for point mutations and deletion or insertion mutations.
- 10 40. The method of Claims 24-26 wherein the expression products are protein molecules.
 - 41. The methods of Claims 24-26 wherein the loss of wild-type GTBP is detected by immunoblotting, e.g. Western blotting.
- 15 42. The methods of Claims 24-26 wherein the alteration of wild type GTBP is detected by immunoenzymology and immunocytochemistry.
 - 43. The method of Claims 24-26 wherein the alteration of wild-type *GTBP* is detected by binding interactions between said GTBP protein and a second cellular protein.
 - 44. The method of Claim 43 wherein the second cellular protein is hMSH2.
 - 45. A method for generating transgenic animals carrying mutant *GTBP* alleles.
 - 46. A pharmaceutical composition useful in the treatment of GTBP-dependent diseases comprising a therapeutically effective amount of GTBP in a pharmaceutically acceptable vehicle.
- 47. A method for supplying wild-type *GTBP* gene function to a cell which has altered GTBP, said gene function being lost by virtue of a mutation in a *GTBP* gene comprising:
- introducing full-length or part of *GTBP* gene in

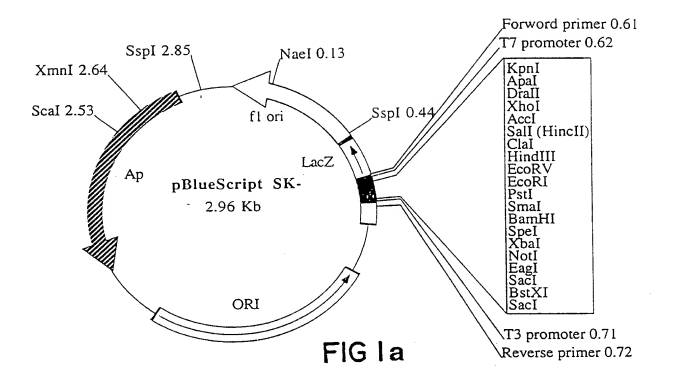
 35 a cell which has lost such gene function such that said
 full-length or part of *GTBP* gene are expressed in the
 cell and encode full-length or part of the GTBP protein

which is capable of complementing the genetic defect at the basis of neoplastic disease.

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48. A method for supplying wild-type GTBP gene function to a cell which has altered GTBP, said gene function being lost by virtue of a mutation in a GTBP gene comprising introducing into a cell a molecule which mimics the effect of GTBP alone or complexed with other molecules.

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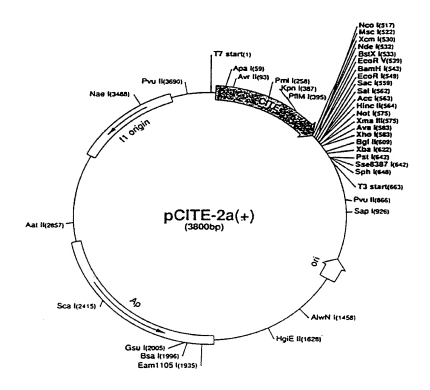
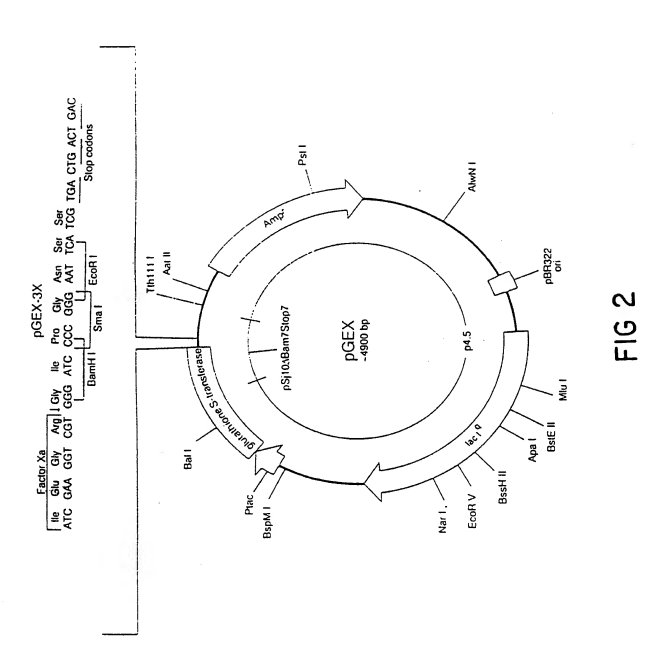
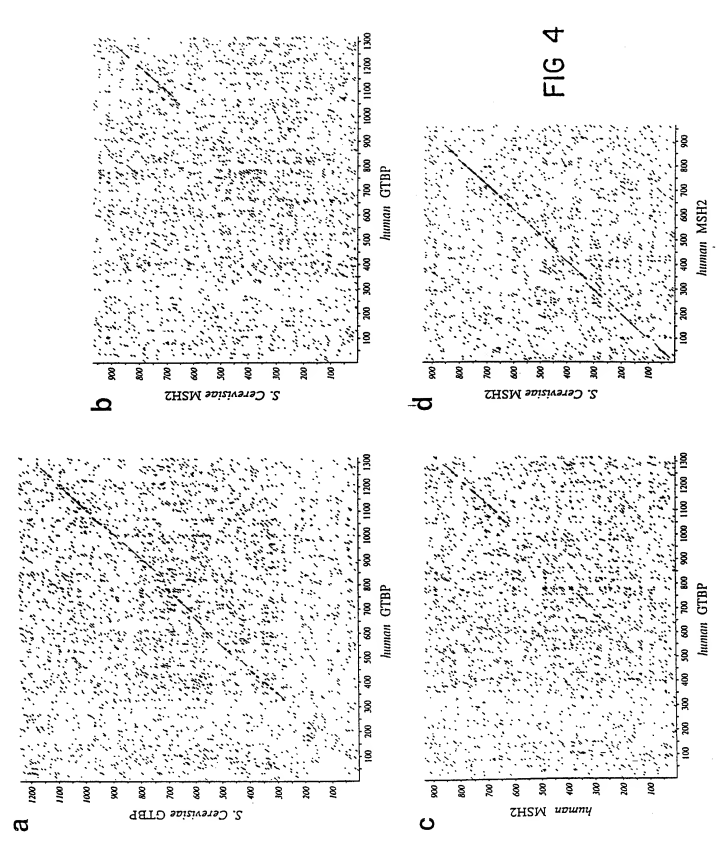


FIG Ib



	3/7
1132	1202
735	805
754	824
680	750
GTBP LATTERNMEGKSTLMRQAGLEAVMAQMGCYVPAEVCRLTPIDRVFTRLGASDRIMSGESTFFVELSETASI	GTBP IMHATAHSLVLVDELGRGTATÄDGTAÄANAVVKELAETIKCRTLFSTHYHSLVEDYSQNVAVRLGHMACM
NMSH2 IITGPNMGGKSTYIROTGVIVLMAQIGCFVPCESAEVSIVDCILARVGAGDSQLKGVSTFMAEWASI	hmsh2 lrsatkdsliidelgrgtstydgfglamaiseyiatrigafcmfrafthfheltalangiptvnnlhvtal
MSH2 IITGPNMGGKSTYIROVGVISLMAQIGCFVPCEEAEIAIVDAILCRVGAGDSQLKGVSTFMVEILETASI	msh2 lrnasknstiivdelgrgtstydgfglamaiaehiaskigcfalfathfheltrelserlpnvknmhvvah
MUTS IITGPNMGGKSTYMRQTALIALMAYIGSYVPAGKVEIDRIFTRVGAADDLASGRSTFMVEMTETANI	muts lhnateyslvlmdeigrgtstydglslamacaenlankiralteathyfeltolpskmegvanvhldal
e z z	P M SM CM



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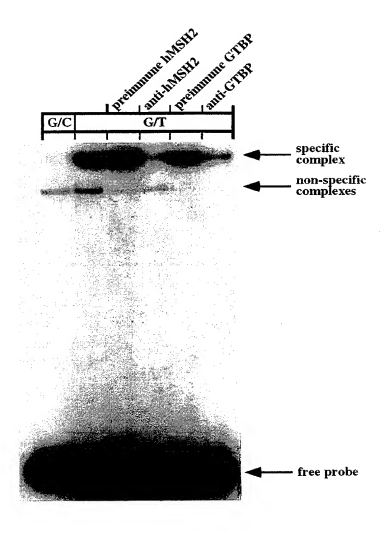


FIG 5

